

L5 ANSWER 3 OF 5 SCISEARCH COPYRIGHT (c) 2007 The Thomson
 Full Text
 Corporation on STN
 Accession Number: 1989:241156 SCISEARCH
 The Genuine Article: U3412
 Title: ONTOGENY OF EPIDERMAL GROWTH-FACTOR RECEPTOR KINASE AND OF LIPOCORTIN-1 IN THE OVINE LUNG
 Author: JOHNSON M D (Reprint); GRAY M E; CARPENTER G; PEPINSKY R B; SUNDELL H; STAHLMAN M T
 Corporate Source: VANDERBILT UNIV, MED CTR, SCH MED, DEPT PEDIAT, NASHVILLE, TN 37232; VANDERBILT UNIV, MED CTR, SCH MED, DEPT BIOCHEM, NASHVILLE, TN 37232; VANDERBILT UNIV, MED CTR, SCH MED, DEPT PATHOL, NASHVILLE, TN 37232; BIOGEN RES CORP, CAMBRIDGE, MA 02142
 Country of Author: USA
 Source: PEDIATRIC RESEARCH, (MAY 1989) Vol. 25, No. 5, pp. 535-541
 ISSN: 0031-3998.
 Publisher: WILLIAMS & WILKINS, 351 WEST CAMDEN ST, BALTIMORE, MD 21201-2436.
 Document Type: Article; Journal
 File Segment: LIFE
 Language: English
 Reference Count: 46
 Entry Date: Entered STN: 1994
 Last Updated on STN: 1994

L5 ANSWER 4 OF 5 SCISEARCH COPYRIGHT (c) 2007 The Thomson
 Full Text
 Corporation on STN
 Accession Number: 1988:48264 SCISEARCH
 The Genuine Article: L7193
 Title: CALCITONIN GENE-RELATED PEPTIDE IN HUMAN-FETAL LUNG AND IN NEONATAL LUNG-DISEASE
 Author: JOHNSON M D (Reprint); GRAY M E; STAHLMAN M T
 Corporate Source: VANDERBILT UNIV, MED CTR, SCH MED, DEPT PATHOL, NASHVILLE, TN 37232; VANDERBILT UNIV, MED CTR, SCH MED, DEPT PEDIAT, NASHVILLE, TN 37232
 Country of Author: USA
 Source: JOURNAL OF HISTOCHEMISTRY & CYTOCHEMISTRY, (FEB 1988) Vol. 36, No. 2, pp. 199-204.
 ISSN: 0022-1954.
 Publisher: HISTOCHEMICAL SOC INC, MT SINAI MEDICAL CENTER 19 EAST 88TH ST SUITE 9G, NEW YORK, NY 10029.
 Document Type: Article; Journal
 File Segment: LIFE
 Language: English
 Reference Count: 2
 Entry Date: Entered STN: 1994
 Last Updated on STN: 1994

L5 ANSWER 5 OF 5 SCISEARCH COPYRIGHT (c) 2007 The Thomson
 Full Text
 Corporation on STN
 Accession Number: 1987:208054 SCISEARCH
 The Genuine Article: G7005
 Title: ONTOGENY OF CALCITONIN-GENE-RELATED PEPTIDE (CGRP) IN HUMAN-FETAL LUNG
 Author: JOHNSON M D (Reprint); GRAY M E; STAHLMAN M T
 Corporate Source: VANDERBILT UNIV, MED CTR, SCH MED, DEPT PEDIAT, NASHVILLE, TN 37232; VANDERBILT UNIV, MED CTR, SCH MED, DEPT PATHOL, NASHVILLE, TN 37232
 Country of Author: USA
 Source: PEDIATRIC RESEARCH, (APR 1987) Vol. 21, No. 4, Part 2, pp. A456-A456.
 ISSN: 0031-3998.
 Publisher: WILLIAMS & WILKINS, 351 WEST CAMDEN ST, BALTIMORE, MD 21201-2436.
 Document Type: Conference; Journal
 File Segment: LIFE

LANGUAGE: English
 REFERENCE COUNT: Entered STN: 1994
 ENTRY DATE: Last Updated on STN: 1994

=> sel L5 4 CIT
 E1 THROUGH E1 ASSIGNED
 => s E1: file CAPLUS: s E1
 L6 21 ("JOHNSON M D, 1988, V36, P199, ?"/RE
 ("JOHNSON M D, 1988, V36, P199, ?"/RE)

FILE 'CAPLUS' ENTERED AT 20:29:54 ON 12 APR 2007
 L7 11 "JOHNSON M D, 1988, V36, P199, ?"/RE
 ("JOHNSON M, 1988, V36, P199, ?"/RE)

=> s 16 or 17
 11 "JOHNSON M D, 1988, V36, P199, ?"/RE
 ("JOHNSON M, 1988, V36, P199, ?"/RE)

L8 11 L6 OR L7

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L6 ANSWER 1 OF 21 SCISEARCH COPYRIGHT (c) 2007 The Thomson
 Full Text
 Corporation on STN
 Accession Number: 2006:803192 SCISEARCH
 The Genuine Article: 073AA
 Title: Serotonin, CGRP, calcitonin, CCK, somatostatin and VIP in the endocrine cells of developing rat lung
 Author: Bayraktar A (Reprint); Tarakci B G
 Corporate Source: Firat Univ, Fac Vet Med, Dept Histol & Embriol, TR-23119

Elazig, Turkey (Reprint)
 Alpayrak@fetal.edu.tr; hkarakis@fetal.edu.tr
 REVUE DE MEDICINE VETERINAIRE, (JUN 2006) Vol. 157, No. 6,
 pp 313-316, 455
 ECOLE NATIONALE VETERINAIRE TOULOUSE, 23 CHEMIN DES
 CARRILES, 31076 TOULOUSE CEDEX 3, FRANCE.
 Article; Journal
 English
 42
 Entered STN: 31 Aug 2006
 Last Updated on STN: 31 Aug 2006
 ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.
 AB Immunoreactivity of serotonin and some regulatory peptides
 (calcitonin gene related peptide (CGRP), calcitonin, cholecystokinin
 (CCK), somatostatin) has been demonstrated in the endocrine cells of
 developing lung by the peroxidase anti-peroxidase method in rat.
 Immunocytochemistry revealed higher density of pulmonary endocrine cells
 containing serotonin and CGRP in foetal and early neonatal periods than in
 the lungs of older rats. Serotonin positive cells were mainly located
 within the bronchial epithelium and in alveolar sacs, whereas the
 localization of CGRP positive cells was essentially restricted to alveolar
 sacs. The calcitonin and somatostatin-containing cells were scarcely
 observed whatever the developmental stages examined in bronchi,
 bronchioles and in alveolar sacs. The CCK expression was weak and
 exclusively found in alveolar sacs, and remained constant from foetus to
 adult stages. VIP immunoreactivity was never detected during lung
 development in rat. These results suggest that serotonin and CGRP would
 be potent mediators involved in the lung ontogeny and in neonatal
 adaptation to air breathing.

L6 ANSWER 2 OF 21 SCISEARCH COPYRIGHT (c) 2007 The Thomson
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 Corporation on STN
 ACCESSION NUMBER: 2004:496303 SCISEARCH
 THE GENUINE ARTICLE: 823VS
 TITLE:
 Forkhead box A2 transcription factor is expressed in all
 types of neuroendocrine lung tumors
 AUTHOR:
 Khoo A (Reprint); Stahlman M T; Johnson J M; Olson S J;
 Whitsett J A
 CORPORATE SOURCE:
 Mayo Clin, Dept Lab Med & Pathol, 4500 San Pablo Rd,
 Jacksonville, FL 32224 USA (Reprint); Mayo Clin, Dept Lab
 Med & Pathol, Jacksonville, FL 32224 USA; Vanderbilt Univ,
 Sch Med, Dept Pathol, Nashville, TN USA; Vanderbilt Univ,
 Sch Med, Dept Pathol, Nashville, TN USA; Univ S Florida,
 Dept Pathol, Coll Med, Tampa, FL USA; James A Haley VA
 Hosp, Tampa, FL USA; Cincinnati Childrens Hosp, Med Ctr,
 Div Neonatol, Cincinnati, OH USA; Cincinnati Childrens
 Hosp, Med Ctr, Div Pulm Biol, Cincinnati, OH USA
 HUMAN PATHOLOGY, (MAY 2004) Vol. 35, No. 5, pp. 560-564.
 ISSN: 0046-8177
 W B SAUNDERS CO, INDEPENDENCE SQUARE WEST CURTIS CENTER,
 STE 300, PHILADELPHIA, PA 19106-3399 USA.
 Article; Journal
 English
 19
 Entered STN: 18 Jun 2004
 Last Updated on STN: 18 Jun 2004
 ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.
 AB Forkhead box A2 (Foxa2) is a winged helix nuclear transcription
 protein that regulates the expression of genes that are critical to lung
 morphogenesis, differentiation, and function, including thyroid
 transcription factor-1, surfactant proteins, and Clara cell secretory
 protein. We examined the immunoreactivity of Foxa2 in paraffin sections
 of 75 lung tumors: 17 typical carcinomas, 2 atypical carcinomas, 4 large
 cell neuroendocrine (NE) carcinomas, 23 small cell carcinomas, 19
 adenocarcinomas, 7 squamous cell carcinomas, and 3 (non-NE) large cell
 carcinomas, using a polyclonal rabbit Foxa2 antibody and a
 biotin-streptavidin detection system. In the adjacent lung, Foxa2 was
 detected in normal and hyperplastic type II cells. Foxa2 immunoreactivity

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 Corporation on STN
 ACCESSION NUMBER: 2002:24655 SCISEARCH
 THE GENUINE ARTICLE: 505JP
 TITLE:
 Hyperplasia of alveolar neuroendocrine cells in rat lung
 carcinogenesis by silica with selective expression of
 proadrenomedullin-derived peptides and amidating enzymes
 AUTHOR:
 Elizegi E, Pino I; Vicent S; Blanco D; Saffioti U;
 Montuenga L M (Reprint)
 CORPORATE SOURCE:
 Univ Navarra, Dept Histol & Pathol, Edif Invest, C
 Irunlarrea 1, Navarra 31008, Spain (Reprint); Univ
 Navarra, Dept Histol & Pathol, Navarra 31008, Spain; NCI,
 Bethesda, MD 20892 USA
 COUNTRY OF AUTHOR:
 Spain; USA
 SOURCE:
 LABORATORY INVESTIGATION, (DEC 2001) Vol. 81, No. 12, pp.
 1627-1638.
 ISSN: 0023-6837
 LIPPINCOTT WILLIAMS & WILKINS, 530 WALNUT ST.
 PHILADELPHIA, PA 19106-3621 USA.
 DOCUMENT TYPE:
 Article; Journal
 LANGUAGE:
 English
 55
 REFERENCE COUNT:
 55
 ENTRY DATE:
 Entered STN: 11 Jan 2002
 Last Updated on STN: 11 Jan 2002
 ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.
 AB Pulmonary neuroendocrine (NE) cells are found as clusters called
 neuroepithelial bodies (NEBs) or as single cells scattered in the
 respiratory epithelium. They express a variety of bioactive peptides, and
 they are thought to be the origin of NE lung tumors. Proadrenomedullin
 N-terminal 20 peptide (PAMP) is a peptide derived from the same precursor
 as adrenomedullin (AM). AM and PAMP are C-terminally amidated during
 their processing by a well-characterized amidating enzyme, peptidylglycine
 alpha-amidating monooxygenase (PAM). We explored AM, PAMP, and PAM
 expression as markers for NE hyperplasia in three rodent species (Fischer
 344 rats, Syrian golden hamsters, and A/J mice) after a single
 intratracheal instillation of crystalline silica (quartz), which was
 previously found to induce different reactions in the three species. Rats
 developed a marked silicosis, with alveolar and bronchiolar hyperplasia
 and formation of peripheral lung epithelial tumors. Mice developed a
 moderate degree of silicosis, but not epithelial hyperplasia or tumors.
 Hamsters showed no silicosis, but not silicosis-related lung NE.
 Cells were immunolabeled for calcitonin gene-related peptide (CGRP), AM,
 PAMP, and PAM in serial sections of lung. The number of positive
 NEBs per lung area and positive cells per NEB were quantified in
 hyperplastic reaction in the NEBs of silica-treated rats occurred only in
 alveolar NEBs, but not in bronchiolar NEBs. From Month 11 onwards, there
 were marked differences in the number of alveolar NEBs per section and in
 the number of cells per alveolar NEB immunoreactive for CGRP. No
 hyperplastic NE cell reaction was observed in silica-treated mice and
 hamsters. Significant PAMP and PAM expression was seen only in rat
 hyperplastic alveolar and in bronchiolar NEBs from Month 11 onwards. In
 E18, rat fetal lung NEBs were found to be strongly positive for PAMP and
 PAM.

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 Corporation on STN
 ACCESSION NUMBER: 2000:823726 SCISEARCH
 THE GENUINE ARTICLE: 367TV
 TITLE:
 Differentiation and proliferation of pulmonary
 neuroendocrine cells
 AUTHOR:
 Ito T (Reprint)

was detected in 13 typical carcinoids (76%), 2 atypical carcinoids (100%),
 2 large cell NE carcinomas (50%), 11 small cell carcinomas (48%), and 1
 adenocarcinoma (5%). Squamous cell carcinomas and (non-NE) large cell
 carcinomas uniformly lacked Foxa2 staining. Expression of Foxa2 in the
 entire spectrum of NE lung tumors is another indication of differentiation
 shared by typical carcinoid, atypical carcinoid, large cell NE carcinoma,
 and small cell carcinoma. Hum PATHOL 35:560-564. (C) 2004 Elsevier Inc.
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L6 ANSWER 3 OF 21 SCISEARCH COPYRIGHT (c) 2007 The Thomson
 Full Text
 Corporation on STN
 ACCESSION NUMBER: 2002:24655 SCISEARCH
 THE GENUINE ARTICLE: 505JP
 TITLE:
 Hyperplasia of alveolar neuroendocrine cells in rat lung
 carcinogenesis by silica with selective expression of
 proadrenomedullin-derived peptides and amidating enzymes
 AUTHOR:
 Elizegi E, Pino I; Vicent S; Blanco D; Saffioti U;
 Montuenga L M (Reprint)
 CORPORATE SOURCE:
 Univ Navarra, Dept Histol & Pathol, Edif Invest, C
 Irunlarrea 1, Navarra 31008, Spain (Reprint); Univ
 Navarra, Dept Histol & Pathol, Navarra 31008, Spain; NCI,
 Bethesda, MD 20892 USA
 COUNTRY OF AUTHOR:
 Spain; USA
 SOURCE:
 LABORATORY INVESTIGATION, (DEC 2001) Vol. 81, No. 12, pp.
 1627-1638.
 ISSN: 0023-6837
 LIPPINCOTT WILLIAMS & WILKINS, 530 WALNUT ST.
 PHILADELPHIA, PA 19106-3621 USA.
 DOCUMENT TYPE:
 Article; Journal
 LANGUAGE:
 English
 55
 REFERENCE COUNT:
 55
 ENTRY DATE:
 Entered STN: 11 Jan 2002
 Last Updated on STN: 11 Jan 2002
 ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.
 AB Pulmonary neuroendocrine (NE) cells are found as clusters called
 neuroepithelial bodies (NEBs) or as single cells scattered in the
 respiratory epithelium. They express a variety of bioactive peptides, and
 they are thought to be the origin of NE lung tumors. Proadrenomedullin
 N-terminal 20 peptide (PAMP) is a peptide derived from the same precursor
 as adrenomedullin (AM). AM and PAMP are C-terminally amidated during
 their processing by a well-characterized amidating enzyme, peptidylglycine
 alpha-amidating monooxygenase (PAM). We explored AM, PAMP, and PAM
 expression as markers for NE hyperplasia in three rodent species (Fischer
 344 rats, Syrian golden hamsters, and A/J mice) after a single
 intratracheal instillation of crystalline silica (quartz), which was
 previously found to induce different reactions in the three species. Rats
 developed a marked silicosis, with alveolar and bronchiolar hyperplasia
 and formation of peripheral lung epithelial tumors. Mice developed a
 moderate degree of silicosis, but not epithelial hyperplasia or tumors.
 Hamsters showed no silicosis, but not silicosis-related lung NE.
 Cells were immunolabeled for calcitonin gene-related peptide (CGRP), AM,
 PAMP, and PAM in serial sections of lung. The number of positive
 NEBs per lung area and positive cells per NEB were quantified in
 hyperplastic reaction in the NEBs of silica-treated rats occurred only in
 alveolar NEBs, but not in bronchiolar NEBs. From Month 11 onwards, there
 were marked differences in the number of alveolar NEBs per section and in
 the number of cells per alveolar NEB immunoreactive for CGRP. No
 hyperplastic NE cell reaction was observed in silica-treated mice and
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 PAM.

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 Corporation on STN
 ACCESSION NUMBER: 2000:823726 SCISEARCH
 THE GENUINE ARTICLE: 367TV
 TITLE:
 Differentiation and proliferation of pulmonary
 neuroendocrine cells
 AUTHOR:
 Ito T (Reprint)

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 Corporation on STN
 ACCESSION NUMBER: 2000:823726 SCISEARCH
 THE GENUINE ARTICLE: 367TV
 TITLE:
 Differentiation and proliferation of pulmonary
 neuroendocrine cells
 AUTHOR:
 Ito T (Reprint)

CORPORATE SOURCE: Yokohama City Univ, Sch Med, Dept Pathol, Kanazawa Ku, 3-9 Fuku Ura, Yokohama, Kanagawa 2360004, Japan (Reprint); Yokohama City Univ, Sch Med, Dept Pathol, Kanazawa Ku, Yokohama, Kanagawa 2360004, Japan

COUNTRY OF AUTHOR: JAPAN

SOURCE: PROGRESS IN HISTOCHEMISTRY AND CYTOCHEMISTRY, (1999) Vol. 34, No. 4, pp. 253-+. ISSN: 0079-6536

PUBLISHER: URBAN & FISCHER VERLAG, GERMAN OFFICE JENA, P O BOX 100537, D-07705 JENA, GERMANY.

DOCUMENT TYPE: General Review; Journal

LANGUAGE: English

REFERENCE COUNT: 356

ENTRY DATE: Entered STN: 2000

ENTRANCE EXAM FOR MD, 2000

• ASPECT IS AVAILABLE IN THE ALL AND IALL FORMATS.

In this review article the morphological profiles of pulmonary neuroendocrine cells (PNEC) in experimental animals and humans are described. Although the mechanisms of differentiation and proliferation of neuroendocrine cells in the airway epithelium remain to be solved, several experimental studies using explant culture and cell culture systems of fetal animal lungs have been performed to clarify fundamental phenomena associated with neuroendocrine differentiation and proliferation. Experimental animal studies using chronic hypoxia, toxic substances, and carcinogens have succeeded in inducing alterations in PNEC systems, and these studies have elucidated the reactions of PNEC in cell injury and inflammation, and functional aspects of PNEC in disease conditions. Human pulmonary neuroendocrine tumors include various histological subtypes, and show divergent morphological and biological varieties. Molecular abnormalities of small cell carcinoma, the most aggressive subtype of pulmonary neuroendocrine tumors, have been extensively studied, but the mechanism of neuroendocrine differentiation of this tumor is still largely unknown.

PNEC share common phenotypes with neuronal cells, and developmental studies have begun contributed evidence that similar transcriptional factors, including the differentially expressed homeodomain (BDH) factors, function in the differentiation of both PNEC and neuronal cells. Such a BDH network may also play a central role in determining cell differentiation in lung carcinomas. Further studies of the neuronal bHLH network, its regulatory system and related signal transduction pathways, will be required for understanding the mechanisms of neuroendocrine differentiation and proliferation in normal and pathological lung conditions.

266 ANSWER 5 OF 21 SCISEARCH COPYRIGHT (c) 2007 The Thomson
 Full Text Corporation on STN
 ACCESSION NUMBER: 1998-602523 SCISEARCH
 THE GENUINE ARTICLE: 108MT necrosis factor induces neuroendocrine
 differentiation in small cell lung cancer cell lines
 TITLE: Haler K J; Patidar K; Zhang F; Emanuel R L; Sunday M E
 AUTHOR: (Reprint)
 CORPORATE SOURCE: Brigham & Women's Hosp, Dept Pathol, 75 Francis St, Boston, MA 02115 USA (Reprint); Brigham & Women's Hosp, Dept Pathol, Boston, MA 02115 USA; Brigham & Women's Hosp, Dept Med, Div Pulm & Crit Care, Boston, MA 02115 USA; Harvard Univ, Sch Med, Boston, MA USA; Childrens Hosp, Dept Pathol, Boston, MA 02115 USA
 COUNTRY OF AUTHOR: USA
 SOURCE: AMERICAN JOURNAL OF PHYSIOLOGY-LUNG CELLULAR AND MOLECULAR PHYSIOLOGY [AUG 1998] Vol. 275, No. 2, pp. L311-L321.
 ISSN: 1040-0605
 PUBLISHER: AMER PHYSIOLOGICAL SOC, 9650 ROCKVILLE PIKE, BETHESDA, MD 20814 USA
 DOCUMENT TYPE: Article; Journal
 LANGUAGE: English
 REFERENCE COUNT: 77
 ENTRY DATE: Entered STN: 1998
 Last Updated on STN: 1998
 ABSTRACT IS AVAILABLE IN THE ALL AND TALL FORMATS
 tumor necrosis factor (TNF)-alpha as a candidate cytokine
 We studied

To promote neuroendocrine cell differentiation in a nitroamine-hyperoxia chamber, rat bronchial epithelial cells were grown in the presence of a 10% serum and/or 10% nitroamine-hyperoxia. Differential screening identified 15 genes that were up-regulated by TNF- α in the presence of nitroamine-hyperoxia. These genes included *CD44*, *CD44v6*, *CD44v7*, *CD44v8*, *CD44v9*, *CD44v10*, *CD44v11*, *CD44v12*, *CD44v13*, *CD44v14*, *CD44v15*, *CD44v16*, *CD44v17*, *CD44v18*, *CD44v19*, *CD44v20*, *CD44v21*, *CD44v22*, *CD44v23*, *CD44v24*, *CD44v25*, *CD44v26*, *CD44v27*, *CD44v28*, *CD44v29*, *CD44v30*, *CD44v31*, *CD44v32*, *CD44v33*, *CD44v34*, *CD44v35*, *CD44v36*, *CD44v37*, *CD44v38*, *CD44v39*, *CD44v40*, *CD44v41*, *CD44v42*, *CD44v43*, *CD44v44*, *CD44v45*, *CD44v46*, *CD44v47*, *CD44v48*, *CD44v49*, *CD44v50*, *CD44v51*, *CD44v52*, *CD44v53*, *CD44v54*, *CD44v55*, *CD44v56*, *CD44v57*, *CD44v58*, *CD44v59*, *CD44v60*, *CD44v61*, *CD44v62*, *CD44v63*, *CD44v64*, *CD44v65*, *CD44v66*, *CD44v67*, *CD44v68*, *CD44v69*, *CD44v70*, *CD44v71*, *CD44v72*, *CD44v73*, *CD44v74*, *CD44v75*, *CD44v76*, *CD44v77*, *CD44v78*, *CD44v79*, *CD44v80*, *CD44v81*, *CD44v82*, *CD44v83*, *CD44v84*, *CD44v85*, *CD44v86*, *CD44v87*, *CD44v88*, *CD44v89*, *CD44v90*, *CD44v91*, *CD44v92*, *CD44v93*, *CD44v94*, *CD44v95*, *CD44v96*, *CD44v97*, *CD44v98*, *CD44v99*, *CD44v100*, *CD44v101*, *CD44v102*, *CD44v103*, *CD44v104*, *CD44v105*, *CD44v106*, *CD44v107*, *CD44v108*, *CD44v109*, *CD44v110*, *CD44v111*, *CD44v112*, *CD44v113*, *CD44v114*, *CD44v115*, *CD44v116*, *CD44v117*, *CD44v118*, *CD44v119*, *CD44v120*, *CD44v121*, *CD44v122*, *CD44v123*, *CD44v124*, *CD44v125*, *CD44v126*, *CD44v127*, *CD44v128*, *CD44v129*, *CD44v130*, *CD44v131*, *CD44v132*, *CD44v133*, *CD44v134*, *CD44v135*, *CD44v136*, *CD44v137*, *CD44v138*, *CD44v139*, *CD44v140*, *CD44v141*, *CD44v142*, *CD44v143*, *CD44v144*, *CD44v145*, *CD44v146*, *CD44v147*, *CD44v148*, *CD44v149*, *CD44v150*, *CD44v151*, *CD44v152*, *CD44v153*, *CD44v154*, *CD44v155*, *CD44v156*, *CD44v157*, *CD44v158*, *CD44v159*, *CD44v160*, *CD44v161*, *CD44v162*, *CD44v163*, *CD44v164*, *CD44v165*, *CD44v166*, *CD44v167*, *CD44v168*, *CD44v169*, *CD44v170*, *CD44v171*, *CD44v172*, *CD44v173*, *CD44v174*, *CD44v175*, *CD44v176*, *CD44v177*, *CD44v178*, *CD44v179*, *CD44v180*, *CD44v181*, *CD44v182*, *CD44v183*, *CD44v184*, *CD44v185*, *CD44v186*, *CD44v187*, *CD44v188*, *CD44v189*, *CD44v190*, *CD44v191*, *CD44v192*, *CD44v193*, *CD44v194*, *CD44v195*, *CD44v196*, *CD44v197*, *CD44v198*, *CD44v199*, *CD44v200*, *CD44v201*, *CD44v202*, *CD44v203*, *CD44v204*, *CD44v205*, *CD44v206*, *CD44v207*, *CD44v208*, *CD44v209*, *CD44v210*, *CD44v211*, *CD44v212*, *CD44v213*, *CD44v214*, *CD44v215*, *CD44v216*, *CD44v217*, *CD44v218*, *CD44v219*, *CD44v220*, *CD44v221*, *CD44v222*, *CD44v223*, *CD44v224*, *CD44v225*, *CD44v226*, *CD44v227*, *CD44v228*, *CD44v229*, *CD44v230*, *CD44v231*, *CD44v232*, *CD44v233*, *CD44v234*, *CD44v235*, *CD44v236*, *CD44v237*, *CD44v238*, *CD44v239*, *CD44v240*, *CD44v241*, *CD44v242*, *CD44v243*, *CD44v244*, *CD44v245*, *CD44v246*, *CD44v247*, *CD44v248*, *CD44v249*, *CD44v250*, *CD44v251*, *CD44v252*, *CD44v253*, *CD44v254*, *CD44v255*, *CD44v256*, *CD44v257*, *CD44v258*, *CD44v259*, *CD44v260*, *CD44v261*, *CD44v262*, *CD44v263*, *CD44v264*, *CD44v265*, *CD44v266*, *CD44v267*, *CD44v268*, *CD44v269*, *CD44v270*, *CD44v271*, *CD44v272*, *CD44v273*, *CD44v274*, *CD44v275*, *CD44v276*, *CD44v277*, *CD44v278*, *CD44v279*, *CD44v280*, *CD44v281*, *CD44v282*, *CD44v283*, *CD44v284*, *CD44v285*, *CD44v286*, *CD44v287*, *CD44v288*, *CD44v289*, *CD44v290*, *CD44v291*, *CD44v292*, *CD44v293*, *CD44v294*, *CD44v295*, *CD44v296*, *CD44v297*, *CD44v298*, *CD44v299*, *CD44v300*, *CD44v301*, *CD44v302*, *CD44v303*, *CD44v304*, *CD44v305*, *CD44v306*, *CD44v307*, *CD44v308*, *CD44v309*, *CD44v310*, *CD44v311*, *CD44v312*, *CD44v313*, *CD44v314*, *CD44v315*, *CD44v316*, *CD44v317*, *CD44v318*, *CD44v319*, *CD44v320*,

66 ANSWER 6 OF 21 SCISEARCH COPYRIGHT (c) 2007 The Thomson
 Full text available on STN
 accession number: 1987-550319 SCISEARCH
 THE GENUINE ARTICLE: XL670
 TITLE: Immunoreactivity for the alpha-subunit of the pituitary
 glycoprotein hormones in pulmonary neuroendocrine cells of
 developing human lung and various perinatal diseases
 VandenSteen P (Reprint); Verbeken E K; VanLommel A;
 Lauweryns J M
 CORPORATE SOURCE: KATHOLIEKE UNIV LEUVEN, SCH MED, LAB HISTOPATHOL, B-3001
 LOUVAIN, BELGIUM
 COUNTRY OF AUTHOR: BELGIUM
 SOURCE: REGULATORY PEPTIDES, (14 MAY 1997) Vol. 70, No. 1, pp.

PUBLISHER: ELSEVIER SCIENCE BV, PO BOX 211, 1000 AE AMSTERDAM, NETHERLANDS.
 DOCUMENT TYPE: Article; Journal
 FILE SEGMENT: LIFE
 LANGUAGE: English
 REFERENCE COUNT: 56
 ENTRY DATE: Entered STN: 1997

ABSTRACT IS AVAILABLE IN THE ALL AND TALL FORMATS*

Infant lung tissue, obtained at autopsy, was studied by immunohistochemistry for the presence of pituitary glycoprotein hormones (PGHs) in the lung. The infants, born at term or preterm, died of various causes. The results provide the first immunological evidence of the presence of the common α -subunit of the pituitary glycoprotein hormones (alpha PGH) in the lung. The immunoreactivity is located in the pulmonary neuroendocrine cells and neuroepithelial bodies. In addition, the cells labelled by alpha PGH antisera (alpha PGH cells) form a subpopulation of the neuroendocrine cells detected by anti-calcitonin immunohistochemistry (CT cells). Moreover, the number of alpha PGH cells appears to increase after neonatal pneumonia or when the number of CT cells is elevated following the development of disease. Also, the weak staining of one of the monoclonal antibodies against the specific β -subunit of thyrotropin (TSH) might, in combination with the increased detectability of α -subunits, indicate that TSH can be endogenously produced in the lung.

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16 ANSER 7 OF 21 SCISEARCH COPYRIGHT (c) 2007 The Thomson
 Full Text
 STN Corporation on
 ACCESSION NUMBER: 1997:478492 SCISEARCH
 THE GENUINE ARTICLE: XF608
 TITLE: Caltonin gene-related peptide immunoreactivity in adult
 mouse lung
 AUTHOR: Verastegui C (Reprint); Oliveira A P; FernandezVivero J;
 Verastegui C (Reprint); Vega J
 INSTITUTION: UNIV CALIF, PAC MED DEPT MORPHOL SCI, CADIZ, SPAIN
 (Reprint)
 CORPORATE SOURCE:

COUNTRY OF AUTHOR: SPAIN
SOURCE: EUROPEAN JOURNAL OF HISTOCHEMISTRY, (1997) Vol. 41, No. 2, 119-126
ISSN: 1121-760X
PUBLISHER: LUIGI PONZIO E FIGLIO, VIA D DA CATALOGNA 1/3, 27100 PAVIA, ITALY
DOCUMENT TYPE: Article; Journal
FILE SEGMENT: LIFE
LANGUAGE: English
REFERENCE COUNT: 41
ENTRY DATE: Entered STN: 1997
Last Updated on STN: 1997

AB Calcitonin gene-related peptide (CGRP) is a 37 amino acid peptide coded by the calcitonin gene that is produced by thyroid C cells and medullary carcinoma. It is also widely distributed in neurons and endocrine cells throughout the body. The presence of CGRP in the lungs suggests that this peptide exerts important regulatory actions at this level, and it can act like a neuroregulator released both from nerve terminals and neuroendocrine (NE) cells. To understand the role of CGRP in the lung, it is important to explore its localization in different species. In this paper, we analyse the presence and localization of CGRP in the adult mouse lung using an immunocytochemical staining method. Our results show a widespread distribution of this peptide in isolated neuroendocrine cells and neuroepithelial bodies (NEBs), as well as in nerve fibres distributed in many areas of the lung, including bronchi and bronchioli. These fibres are in close contact with epithelium, neuroendocrine cells and smooth muscle. In addition, some immunostained nerve cell bodies and immunoreactive intrinsic ganglion cells can be shown. CGRP has been previously demonstrated in the mammalian lung using immunocytochemistry. To the best of our knowledge, this is the first time that CGRP has been immunocytochemically demonstrated in the mouse lung both in NE cells, NEBs, ganglion cells and in nerve fibres which are related to neuroendocrine cells.

L6 ANSWER 8 OF 21 SCISEARCH COPYRIGHT (c) 2007 The Thomson
Full Text
Accession on STN
ACCESSION NUMBER: 1997:355866 SCISEARCH
THE GENUINE ARTICLE: W4198
TITLE: Quantitative microscopical methods for the identification and localisation of nerves and neuroendocrine cell markers in mammalian lung
AUTHOR: SOROKIN S P (Reprint); Hoyt R F; Shaffer M J
CORPORATE SOURCE: BOSTON UNIV, SCH MED, DEPT ANAT & NEUROBIOL, 80 E CONCORD ST, BOSTON, MA 02118 (Reprint)
COUNTRY OF AUTHOR: USA
SOURCE: MICROSCOPY RESEARCH AND TECHNIQUE, (1 APR 1997) Vol. 37, No. 1, pp. 43-61.
ISSN: 1059-910X
PUBLISHER: WILEY-LISS, DIV JOHN WILEY & SONS INC, 605 THIRD AVE, NEW YORK, NY 10158-0012.
DOCUMENT TYPE: General Review; Journal
FILE SEGMENT: LIFE; ENGI
LANGUAGE: English
REFERENCE COUNT: 114
ENTRY DATE: Entered STN: 1997
Last Updated on STN: 1997

AB This paper summarizes current knowledge and advances speculation about the formation of the neuroendocrine system of mammalian lungs (comprising uninnervated solitary and clustered small-granule cells and innervated neuroepithelial bodies). It relates the initial appearance of neuroendocrine cells to regulation of mitotic activity in the epithelium during the development of the lung and pays special attention to the later ingrowth of nerves that converts some of them into neuroepithelial bodies. Structures considered ideally adapted to function as chemoreceptors. A few original observations from ongoing immunohistochemical, electron microscopic, and analytical studies have been included here and there to point the discussion. The neuroendocrine cells are derived from undifferentiated precursors present in the endodermal pulmonary epithelium. At an early pseudoglandular stage of lung development these precursors begin to differentiate into neuroendocrine small-granule cells, innervating airways and upper trachea, the expanding centrifugally subsequent many of the intrapulmonary small-granule cell clusters become innervated. This event the delayed appearance of small-granule cells synthesizing other than the dominant peptides and amines (calcitonin gene-related peptide and serotonin in rodents, gastrin-releasing peptide and serotonin in human beings), and other regional adjustments yield the population distribution present in the lungs of adults. Neuroendocrine cell precursors normally differentiate into typical serotonin- or peptide-synthesizing small-granule cells without requiring direct contact by nerves, and dissociated cells from a previously innervated population continue to exhibit physiological characteristics of oxygen sensors despite the loss of contact with nerves. Development of the innervation occurs in stages. Small-granule cell clusters are reached first by ganglion cells derived from pulmonary neuroblasts and later on by processes of extrinsic sensory nerves. The latter not only convey information to the central nervous system but also serve in a variety of ways to extend the neuroepithelial bodies' sphere of influence within the lung itself. (C) 1997 Wiley-Liss, Inc.

L6 ANSWER 9 OF 21 SCISEARCH COPYRIGHT (c) 2007 The Thomson
Full Text
Accession on STN
ACCESSION NUMBER: 1997:355862 SCISEARCH
THE GENUINE ARTICLE: W4198
TITLE: Ontogeny of neuroepithelial bodies: Correlations with mitogenesis and innervation
AUTHOR: SOROKIN S P (Reprint); Hoyt R F; Shaffer M J
CORPORATE SOURCE: BOSTON UNIV, SCH MED, DEPT ANAT & NEUROBIOL, 80 E CONCORD ST, BOSTON, MA 02118 (Reprint)
COUNTRY OF AUTHOR: USA
SOURCE: MICROSCOPY RESEARCH AND TECHNIQUE, (1 APR 1997) Vol. 37, No. 1, pp. 43-61.
ISSN: 1059-910X
PUBLISHER: WILEY-LISS, DIV JOHN WILEY & SONS INC, 605 THIRD AVE, NEW YORK, NY 10158-0012.
DOCUMENT TYPE: General Review; Journal
FILE SEGMENT: LIFE; ENGI
LANGUAGE: English
REFERENCE COUNT: 114
ENTRY DATE: Entered STN: 1997
Last Updated on STN: 1997

AB This paper summarizes current knowledge and advances speculation about the formation of the neuroendocrine system of mammalian lungs (comprising uninnervated solitary and clustered small-granule cells and innervated neuroepithelial bodies). It relates the initial appearance of neuroendocrine cells to regulation of mitotic activity in the epithelium during the development of the lung and pays special attention to the later ingrowth of nerves that converts some of them into neuroepithelial bodies. Structures considered ideally adapted to function as chemoreceptors. A few original observations from ongoing immunohistochemical, electron microscopic, and analytical studies have been included here and there to point the discussion. The neuroendocrine cells are derived from undifferentiated precursors present in the endodermal pulmonary epithelium. At an early pseudoglandular stage of lung development these precursors begin to differentiate into neuroendocrine small-granule cells, innervating airways and upper trachea, the expanding centrifugally subsequent many of the intrapulmonary small-granule cell clusters become innervated. This event the delayed appearance of small-granule cells synthesizing other than the dominant peptides and amines (calcitonin gene-related peptide and serotonin in rodents, gastrin-releasing peptide and serotonin in human beings), and other regional adjustments yield the population distribution present in the lungs of adults. Neuroendocrine cell precursors normally differentiate into typical serotonin- or peptide-synthesizing small-granule cells without requiring direct contact by nerves, and dissociated cells from a previously innervated population continue to exhibit physiological characteristics of oxygen sensors despite the loss of contact with nerves. Development of the innervation occurs in stages. Small-granule cell clusters are reached first by ganglion cells derived from pulmonary neuroblasts and later on by processes of extrinsic sensory nerves. The latter not only convey information to the central nervous system but also serve in a variety of ways to extend the neuroepithelial bodies' sphere of influence within the lung itself. (C) 1997 Wiley-Liss, Inc.

L6 ANSWER 10 OF 21 SCISEARCH COPYRIGHT (c) 2007 The Thomson
Full Text
Accession on STN
ACCESSION NUMBER: 1997:19565 SCISEARCH
THE GENUINE ARTICLE: V2426
TITLE: Ontogeny of Clara cell-specific protein and its mRNA:

colocalize immunoreactivity for serotonin (5-HT) and calcitonin gene-related peptide (CGRP) and persist throughout adult life. Postnatally, avian cells also express calcitonin (CT), but associated and CGRP staining correlates with the onset of local, NEB-associated mitogenesis in fetal hamster airway epithelium. The second pattern begins after birth and is unique to the larynx and cartilaginous trachea. It involves differentiation of single cells which stain for CGRP but not 5-HT. Later, a proportion also stain for CT. This pattern seemingly accounts for the predominance of single cells in laryngotracheal epithelium of adult animals. In the third pattern, cells immunoreactive for peptide YY (PYY) differentiate, singly at first and later among cells of tiny pNEBs. This begins postnatally in alveoli, spreading centripetally with retrograde differentiation of alveolar epithelium back into the bronchiolar terminations. Restricted distribution and lack of immunoreactivity for 5-HT, CGRP, or CT suggest that the PYY-positive endocrine cells form a regional subset performing special roles in pulmonary homeostasis.

L6 ANSWER 14 OF 21 SCISEARCH COPYRIGHT (c) 2007 The Thomson

Full Text

ACCESSION NUMBER: 1994:2382 SCISEARCH

FILE SEGMENT: NM222

TITLE: ONTOGENY OF ENDOCRINE-CELLS IN THE RESPIRATORY SYSTEM OF SYRIAN GOLDEN-HAMSTERS .1. LARYNX AND TRACHEA

AUTHOR: SYRIAN GOLDEN-HAMSTERS .1. LARYNX AND TRACHEA

CORPORATE SOURCE: SYRIAN GOLDEN-HAMSTERS .1. LARYNX AND TRACHEA

COUNTRY OF AUTHOR: SYRIAN GOLDEN-HAMSTERS .1. LARYNX AND TRACHEA

SOURCE: SYRIAN GOLDEN-HAMSTERS .1. LARYNX AND TRACHEA

ISSN: 0302-766X

PUBLISHER: SPRINGER VERLAG, 175 FIFTH AVE, NEW YORK, NY 10010.

DOCUMENT TYPE: Article; Journal

FILE SEGMENT: Life

LANGUAGE: English

REFERENCE COUNT: 47

ENTRY DATE: Entered STN: 1994

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS*

The ontogeny of protein gene product 9.5 (PGP 9.5), serotonin (5-HT), calcitonin gene-related peptide (CGRP), and calcitonin (CT) immunoreactivity was evaluated in small-granule endocrine cells of hamster laryngotracheal epithelium from fetal day 11 to adulthood. Two first patterns (beginning during fetal day 11) of differentiation occur. The first pattern begins during fetal day 11. Endoneurial bud-like and clustered colocalize immunoreactivity for PGP 9.5, 5-HT, and CGRP in the larynx and proximal 2/3 of the trachea on day 12 and spread to the caudal trachea on day 13. 5-HT disappears fleetingly during the 24 h preceding birth; otherwise immunoreactivity for all three substances persists into adulthood. The clusters of endocrine cells survive beyond birth but are so diluted by expansion of the nonendocrine epithelium as to become inconspicuous. Since innervation was not actually observed, these clusters may persist as pNEBs, without developing connections to afferent or efferent nerve fibers. The second pattern concerns single small-granule cells stainable for CGRP but not for 5-HT. These cells first appear in the larynx and cartilaginous part of the cranial trachea on postnatal day 3, and in the middle and caudal trachea, on day 5. The cells increase in number on day 7. In adults, they predominate among endocrine cells of the cartilaginous region. A subset of these cells begins to co-express CT proximally on postnatal day 10, reaching the caudal end of the trachea by 3 weeks. A few elements of the older 5-HT-positive population may also become immunoreactive for CT in juvenile hamsters.

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Full Text

ACCESSION NUMBER: 1993:420272 SCISEARCH

FILE SEGMENT: NM222

TITLE: ONTOGENY OF ENDOCRINE-CELLS IN THE RESPIRATORY SYSTEM OF SYRIAN GOLDEN-HAMSTERS .1. LARYNX AND TRACHEA

AUTHOR: SYRIAN GOLDEN-HAMSTERS .1. LARYNX AND TRACHEA

CORPORATE SOURCE: SYRIAN GOLDEN-HAMSTERS .1. LARYNX AND TRACHEA

COUNTRY OF AUTHOR: SYRIAN GOLDEN-HAMSTERS .1. LARYNX AND TRACHEA

SOURCE: SYRIAN GOLDEN-HAMSTERS .1. LARYNX AND TRACHEA

ISSN: 0302-766X

PUBLISHER: SPRINGER VERLAG, 175 FIFTH AVE, NEW YORK, NY 10010.

DOCUMENT TYPE: Article; Journal

FILE SEGMENT: Life

LANGUAGE: English

REFERENCE COUNT: 47

ENTRY DATE: Entered STN: 1994

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS*

The ontogeny of protein gene product 9.5 (PGP 9.5), serotonin (5-HT), calcitonin gene-related peptide (CGRP), and calcitonin (CT) immunoreactivity was evaluated in small-granule endocrine cells of hamster laryngotracheal epithelium from fetal day 11 to adulthood. Two first patterns (beginning during fetal day 11) of differentiation occur. The first pattern begins during fetal day 11. Endoneurial bud-like and clustered colocalize immunoreactivity for PGP 9.5, 5-HT, and CGRP in the larynx and proximal 2/3 of the trachea on day 12 and spread to the caudal trachea on day 13. 5-HT disappears fleetingly during the 24 h preceding birth; otherwise immunoreactivity for all three substances persists into adulthood. The clusters of endocrine cells survive beyond birth but are so diluted by expansion of the nonendocrine epithelium as to become inconspicuous. Since innervation was not actually observed, these clusters may persist as pNEBs, without developing connections to afferent or efferent nerve fibers. The second pattern concerns single small-granule cells stainable for CGRP but not for 5-HT. These cells first appear in the larynx and cartilaginous part of the cranial trachea on postnatal day 3, and in the middle and caudal trachea, on day 5. The cells increase in number on day 7. In adults, they predominate among endocrine cells of the cartilaginous region. A subset of these cells begins to co-express CT proximally on postnatal day 10, reaching the caudal end of the trachea by 3 weeks. A few elements of the older 5-HT-positive population may also become immunoreactive for CT in juvenile hamsters.

TITLE: IMMUNOCYTOCHEMICAL STUDY OF THE LUNG OF DOMESTIC-FOWL AND PIGEON ENDOCRINE CELLS AND NERVES

AUTHOR: LAPEZ J (REPRINT); BARRECHEN A; A: SESMA P

CORPORATE SOURCE: UNIV NAVARRA, E-31080 PAMPLONA, SPAIN

COUNTRY OF AUTHOR: UNIV NAVARRA, E-31080 PAMPLONA, SPAIN

SOURCE: UNIV NAVARRA, E-31080 PAMPLONA, SPAIN

ISSN: 0302-766X

PUBLISHER: SPRINGER VERLAG, 175 FIFTH AVE, NEW YORK, NY 10010.

DOCUMENT TYPE: Article; Journal

FILE SEGMENT: Life

LANGUAGE: English

REFERENCE COUNT: 58

ENTRY DATE: Entered STN: 1994

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS*

The presence of endocrine cells and nerves in the lung of 2 avian species (Gallus gallus and Columba livia domestica) has been studied by peroxidase-antiperoxidase (PAP) and avidin-biotin complex (ABC) immunocytochemical methods at the light-microscopic level. Two immunoreactive cell-types have been identified in the epithelium of the primary and secondary bronchi of chick lung: serotonin- and bombesin-immunoreactive cells; and 3 cell-types, namely, serotonin-, bombesin- and CGRP- (calcitonin gene related peptide) immunoreactive cells. Have been located in the bronchial epithelium of pigeon lung. Co-localization of 2 different immunoreactivities within the same cell has not been detected. VIP-immunoreactive nerves have been observed in different locations in chick lung.

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Full Text

ACCESSION NUMBER: 1993:290825 SCISEARCH

FILE SEGMENT: K2646

TITLE: COLOCALIZATION OF PEPTIDE-HORMONES IN NEUROENDOCRINE CELLS OF HUMAN FETAL AND NEWBORN LUNGS - AN ELECTRON-MICROSCOPIC STUDY

AUTHOR: STAHLMAN M T (REPRINT); GRAY M E

CORPORATE SOURCE: VANDERBILT UNIV, MED CTR, SCH MED, DEPT PEDIAT, A-0126 MED

COUNTRY OF AUTHOR: USA

SOURCE: USA

ISSN: 0003-276X

PUBLISHER: WILEY-LISS, 605 THIRD AVE, NEW YORK, NY 10158-0012.

DOCUMENT TYPE: Article; Journal

FILE SEGMENT: Life

LANGUAGE: English

REFERENCE COUNT: 26

ENTRY DATE: Entered STN: 1994

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS*

This study investigated the colocalization of the peptide hormones bombesin or calcitonin with calcitonin gene related peptide (CGRP) in neuroendocrine cells (NE) in the lungs of human fetuses of varying gestational ages and in the lungs of newborn infants who died with acute or chronic lung disease in the first weeks or months after birth. Double immunolabeling of dense core granules for these peptides was also studied in this same patient population. On-grid double gold immunolabeling was carried out on 29 subjects using anti-bombesin and anti-CGRP and on 22 subjects using anti-calcitonin and anti-CGRP as primary antibodies. The secondary antibodies being labeled with different-size gold spheres. Colocalization of both bombesin and calcitonin with CGRP was demonstrated. not only in the same NE cell, but also on the same dense core granule. Colocalization was rarely found in normal fetuses, and most frequently found in newborn infants with acute lung disease, usually hyaline membrane disease (HMD), or with the development of chronic lung disease in the first weeks or months after birth. Double labeling of the same dense core granules might imply action of peptides in concert, or perhaps one peptide

acting in a paracrine role (e.g., on bronchial or bronchiolar smooth muscle) and the second peptide acting in an autocrine fashion on the parent cell (e.g., in the regulation of granule production or release).

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Full Text
Accession on STN
Accession Number: 1993:290808 SCISEARCH
The Genuine Article: K2646
Title: NEUROENDOCRINE CELLS AND NERVES OF THE LUNG
Author: ADRIANSEN D (Reprint); SCHEUERMANN D W
Corporate Source: UNIV ANTWERP, CELL BIOL & HISTOL, GROENENBORGERLAAN 171, B-2020 ANTWERP, BELGIUM
Country of Author: BELGIUM
Source: ANATOMICAL RECORD, (MAY 1993) Vol. 236, No. 1, pp. 70-86.
ISSN: 0003-276X
Publisher: WILEY-LISS, DIV JOHN WILEY & SONS INC 605 THIRD AVE, NEW YORK, NY 10158-0012.
Document Type: Article; Journal
File Segment: LIFE
Language: English
Reference Count: 168
Entry Date: Entered STN: 1994
Last Updated on STN: 1994

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Full Text
Accession on STN
Accession Number: 1993:92183 SCISEARCH
The Genuine Article: KK751
Title: PULMONARY BLASTOMA - COMPARISON BETWEEN ITS EPITHELIAL COMPONENTS AND FETAL BRONCHIAL EPITHELIUM
Author: INOUE H (Reprint); KASAI K; SHINADA J; YOSHIMURA H; KAMEYA T
Corporate Source: KITASATO UNIV, SCH MED, DEPT PATHOL, KITASATO 1-15-1, SAGAMIHARA, KANAGAWA 228, JAPAN (Reprint); KITASATO UNIV, SCH MED, DEPT THORAC SURG, SAGAMIHARA, KANAGAWA 228, JAPAN
Country of Author: JAPAN
Source: ACTA PATHOLOGICA JAPONICA, (DEC 1992) Vol. 42, No. 12, pp. 884-892.
ISSN: 0001-6632
Publisher: BLACKWELL PUBLISHING ASIA, 54 UNIVERSITY ST, P O BOX 378, CARLTON, VICTORIA 3053, AUSTRALIA.
Document Type: Article; Journal
Language: English
Reference Count: 20
Entry Date: Entered STN: 1994
Last Updated on STN: 1994

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS
AB Three cases of pulmonary blastoma exhibiting biphasic epithelial and stromal patterns, and a case of fetal lung-type adenocarcinoma were examined by immunohistochemistry and electron microscopy (EM) and compared with fetal bronchial epithelium in order to explore the multidirectional differentiation of their epithelial components. The glandular cells of all four tumors resembled fetal bronchial epithelial cells in the pseudoglandular stage. Neuroendocrine (NE) cells were also present; they were argyrophilic and expressed pan-NE markers, neurosecretory granules and peptide hormones. The neural cell adhesion molecule (NCAM) was strongly expressed on the cell membranes of glandular cells, as in the case of proximal bronchial epithelial cells at the pseudoglandular stage in fetal lung. Sialosylated Lewis(x) was also expressed, indicating that the epithelial cells were possibly of endodermal origin. Two of the four cases showed considerable immunoreactivity for alpha-fetoprotein (AFP). The epithelial cells of pulmonary blastomas may occasionally dedifferentiate into cells functionally resembling fetal hepatic, foregut and yolk sac cells expressing AFP. Tumor examination by immunohistochemistry and EM suggested that the glandular cells of the tumors may differentiate to some extent like those of fetal large bronchi at the pseudoglandular stage, but there was concordance and discordance in the expression of neuroendocrine and oncofetal markers between blastomatous tumors and fetal bronchial epithelium.

L6 ANSWER 19 OF 21 SCISEARCH COPYRIGHT (c) 2007 The Thomson
Full Text
Accession on STN
Accession Number: 1992:662720 SCISEARCH
The Genuine Article: JX081
Title: COMPARATIVE HISTOLOGICAL OVERVIEW OF THE CHEMICAL CODING OF THE PULMONARY NEUROEPITHELIAL ENDOCRINE SYSTEM IN HEALTH AND DISEASE
Author: SCHEUERMANN D W (Reprint); ADRIANSEN D; TIMMERMAN J P; DEGRUOTLASSEEL M H A
Corporate Source: UNIV ANTWERP, INST HISTOL & MICROSCOP ANAT, GROENENBORGERLAAN 171, B-2020 ANTWERP, BELGIUM (Reprint)
Country of Author: BELGIUM
Source: EUROPEAN JOURNAL OF MORPHOLOGY, (1992) Vol. 30, No. 2, pp. 101-112.
ISSN: 0924-3860
Publisher: SWETS ZEITLINGER PUBLISHERS, P O BOX 825, 2160 SZ LISSE, NETHERLANDS.
Document Type: General Review; Journal
File Segment: LIFE
Language: English
Reference Count: 80
Entry Date: Entered STN: 1994
Last Updated on STN: 1994

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS
AB Pulmonary neuroepithelial endocrine cells have been shown to contain serotonergic immunoreactivity in almost every species studied. Regulatory peptides, of which at least ten have been reported so far, were mostly only demonstrated in a number of the investigated species or in a subpopulation of neuroepithelial endocrine cells. Calcitonin gene-related peptide, calcitonin, bombesin/gastrin-releasing peptide, enkephalin, somatostatin, substance P, cholecystokinin and polypeptide YY were found in normal lung tissues, whereas ACTH and several other bioactive substances should be regarded as ectopic. The human pulmonary neuroepithelial endocrine system seems to harbour the largest spectrum of bioactive mediators.

The distribution patterns of bioactive substances in various subpopulations of solitary neuroepithelial endocrine cells or neuroepithelial bodies and in different cells of a single neuroepithelial body reveal a great complexity. Therefore, further research is needed to elucidate the chemical coding of this system.

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Full Text
Accession on STN
Accession Number: 1992:46773 SCISEARCH
The Genuine Article: GY785
Title: ISOLATION AND CULTURE OF NEUROENDOCRINE CELLS FROM FETAL RABBIT LUNG USING IMMUNOMAGNETIC TECHNIQUES
Author: SPEIRS V (Reprint); WANG Y Y; YEEGER H; CUTZ E
Corporate Source: HOSP SICK CHILDREN, DEPT PATHOL, 555 UNIV AVE, TORONTO M5G 1X8, ONTARIO, CANADA (Reprint); HOSP SICK CHILDREN RES INST, TORONTO M5G 1X8, ONTARIO, CANADA; UNIV TORONTO, TORONTO M5S 1A1, ONTARIO, CANADA
Country of Author: CANADA
Source: AMERICAN JOURNAL OF RESPIRATORY CELL AND MOLECULAR BIOLOGY (JAN 1992) Vol. 6, No. 1, pp. 63-67.
ISSN: 1044-1549
Publisher: AMER LUNG ASSOC, 1740 BROADWAY, NEW YORK, NY 10019.
Document Type: Article; Journal
File Segment: LIFE
Language: English
Reference Count: 32
Entry Date: Entered STN: 1994
Last Updated on STN: 1994

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS
AB We describe a novel method for the isolation and subsequent culture of pulmonary neuroendocrine cells (PNEC) from normal fetal rabbit lung using immunomagnetic techniques with a monoclonal antibody, MOC-1. This surface antigen has originally been identified on small cell carcinoma of the lung. Our immunohistochemical studies have shown that MOC-1 cross-reacts with PNEC of human and rabbit fetal lungs on frozen sections,

and in fixed cultures of rabbit fetal lung. Using a combination of mechanical and enzymatic disaggregation, single cell suspension of fetal rabbit lung was obtained. These cells were incubated with ECM conjugated to magnetic beads. These cells were selectively released from the heterogeneous mixture using a magnet giving up to 2-fold enrichment compared with our previously reported method. These cells were maintained in culture in a functional state for up to 7 days. The ability to prepare PNEC from rabbit fetal lung offers an opportunity to develop in vitro models to investigate the physiologic and biochemical properties of these cells, and ultimately it may lead to a better understanding of their function in health and disease.

L6 ANSWER 21 OF 21 SCISEARCH COPYRIGHT (c) 2007 The Thomson

Full Text

Accession Number: 1991:65302 SCISEARCH
The Genuine Article: EV449
Title: GASTRIN-RELEASING PEPTIDE GENE-PRODUCTS IN MIDTRIMESTER HUMAN FETAL LUNG WITH AND WITHOUT MATERNAL SMOKING HISTORY DURING PREGNANCY

Author: CHEN M F (Reprint); LEWIS S J; JAGOE R; ALEXANDER N; VANNOORDEN S; SPRINGALL D R; POLAK J M

Corporate Source: MCGILL UNIV, DEPT PATHOL, MONTREAL H3A 2B4, QUEBEC, CANADA; HAMMERSMITH HOSP, ROYAL POSTGRAD MED SCH, DEPT HISTOCHEM, LONDON W12 0HS, ENGLAND; HAMMERSMITH HOSP, ROYAL POSTGRAD MED SCH, DEPT MED PHYS, LONDON W12 0HS, ENGLAND

COUNTRY OF AUTHOR: CANADA; ENGLAND
SOURCE: PEDIATRIC PULMONOLOGY, (1991) Vol. 10, No. 1, PP. 30-35. ISSN: 8755-6863.

PUBLISHER: WILEY-LISS, DIV JOHN WILEY & SONS INC 605 THIRD AVE, NEW YORK, NY 10158-0012.

DOCUMENT TYPE: Article; Journal

FILE SEGMENT: CLIN

LANGUAGE: English

REFERENCE COUNT: 18

ENTRY DATE: Entered STN: 1994

Last Updated on STN: 1994

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AB A preliminary morphological study on human fetal lungs with positive maternal smoking history demonstrated alterations of the neuroepithelial bodies (NEBs). We studied human fetal lung tissue between the gestational ages of 12 weeks and 19 weeks, comprising 12 cases with a smoking history during pregnancy (group 1) and eight cases without a smoking history during pregnancy (group 2). We demonstrated, by immunocytochemistry, the presence of gastrin-releasing peptide (GRP), GRP-C-flanking peptide, in all 20 cases. Identification of the neuroepithelial cells (NECs) made by computer-enhanced image analysis using the Context Vision system expressing 1) the total stained areas of the NECs per unit area of section and 2) the total staining areas of the NECs per unit area of airway epithelium, measured as the area of cytokeratin immunoreactivity in an adjacent section. The results show no statistically significant difference between groups 1 and 2 for either GRP 14-27 or C-flanking peptides. The apparent lack of influence of maternal smoking during pregnancy on the expression of GRP gene products in the NECs could be a reflection of inherently reduced reactivity of the cells during the gestation period studied. However, a larger series is needed before any conclusions can be made. Alternatively, the adverse effects of smoking might be reflected during the canalicular phase of lung development; an increased immunoreactivity appears to be present during that period. The expression of pro-GRP gene products in the pulmonary NECs of older fetuses and neonates with maternal smoking history during pregnancy requires further study.

=> d his

(FILE 'HOME' ENTERED AT 20:25:55 ON 12 APR 2007)

FILE 'SCISEARCH' ENTERED AT 20:26:13 ON 12 APR 2007

L1 198 S STAHLMAN M7/AU

L2 1797 S GRAY M7/AU
L3 149 S L AND L2
L4 11201 S JOHNSON M7/AU
L5 5 L4 AND L3
L6 21 S E1
SEL L5 4 CIT

FILE 'CAPLUS' ENTERED AT 20:29:54 ON 12 APR 2007

L7 11 S E1

L8 11 S L6 OR L7

FILE 'CAPLUS' ENTERED AT 20:32:20 ON 12 APR 2007

FILE 'SCISEARCH' ENTERED AT 20:32:42 ON 12 APR 2007

FILE 'CAPLUS' ENTERED AT 20:32:43 ON 12 APR 2007

=> 17 not L6

L7 IS NOT A RECOGNIZED COMMAND

The previous command name entered was not recognized by the system.

For a list of commands available to you in the current file, enter

"HELP COMMANDS" at an arrow prompt (=>).

=> s 17 not L6

L11 "JOHNSON M D, 1988, V36, P199-7"/RE

L9 0 L7 NOT L6

=> file SCISEARCH; sel L6 11 CIT

FILE 'SCISEARCH' ENTERED AT 20:50:35 ON 12 APR 2007

E2 THROUGH E2 ASSIGNED

=> s E2

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Full Text

Accession Number: 2005:872454 SCISEARCH

The Genuine Article: Fetal oxygen tension promotes Tenascin-C-dependent lung branching morphogenesis

Author: KISHIMOTO, K; VAUGHN, J; MCKEAN, D; JONES, P L
Corporate Source: Univ Colorado, Hlth Sci Ctr, Dept Med, CO 80262 USA
Lab Box B133, 4200 E 9th Ave, Denver, CO 80262 USA
(Reprint): Univ Colorado, Hlth Sci Ctr, Dept Med, Colorado Pulm Res Lab, Denver, CO 80262 USA; Univ Colorado, Hlth Sci Ctr, Dept Pediat, Denver, CO 80202 USA
Country of Author: USA
Source: DEVELOPMENTAL DYNAMICS, (SEP 2005) Vol. 234, No. 1, PP. 1-10. ISSN: 1058-8388.

PUBLISHER: WILEY-LISS, DIV JOHN WILEY & SONS INC, 111 RIVER ST, HOBOKEN, NJ 07030 USA.

DOCUMENT TYPE: Article; Journal

LANGUAGE: English

REFERENCE COUNT: 51

ENTRY DATE: Entered STN: 8 Sep 2005

Last Updated on STN: 8 Sep 2005

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AB Tenascin-C (TN-C) is a mesenchyme-derived extracellular matrix (ECM) glycoprotein required for fetal lung branching morphogenesis. Given that the low oxygen (O-2) environment of the fetus is also essential for normal lung branching morphogenesis, we determined whether fetal O-2 tension supports this process by promoting TN-C expression. Initial studies showed that 15-day fetal rat lung explants cultured for 2 days at 3 & O-2

ACCESSION NUMBER: 1999:241913 SCISEARCH
 THE GENUINE ARTICLE: 182zy
 TITLE: The pulmonary neuroendocrine system: The past decade
 AUTHOR: Van Lommel (Reprint); Ballez; Fannet; W. J. M.
 CORPORATE SOURCE: Katholieke Univ. Leuven, Fac Med, Anat Pathol Lab, B-3000
 Leuven, Belgium
 COUNTRY OF AUTHOR: Belgium
 SOURCE: ARCHIVES OF HISTOLOGY AND CYTOLOGY, (MAR 1999) Vol. 62,
 No. 1, pp. 1-16.
 PUBLISHER: JAPAN SOC HISTOL DOCUMENTATION NIIGATA UNIV MEDICAL SCHOOL
 DEPARTMENT OF ANATOMY ASAHI-MACHI, NIIGATA, 951, JAPAN.
 DOCUMENT TYPE: General Review; Journal
 LANGUAGE: English
 REFERENCE COUNT: 115
 ENTRY DATE: Entered STN: 1999
 Last Updated on STN: 1999

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.
 AB The pulmonary neuroendocrine system consists of specialized airway
 endocrine epithelial cells, associated with nerve fibres. The epithelial
 cells, the pulmonary neuroendocrine cells (PNEC), can be solitary or
 clustered to form neuroendocrine bodies (NEB). During the last thirty
 years, the pulmonary neuroendocrine system has been intensively
 investigated and much knowledge of its function has been obtained. This
 text reviews work which dates from the last ten years. In this period, the
 picture of the pulmonary neuroendocrine system we previously had, has not
 fundamentally changed. The pulmonary neuroendocrine system is still
 regarded as an oxygen sensitive chemoreceptor with local and
 reflex-mediated regulatory functions, and as a regulator of airway growth
 and development. Continuing research has much more refined this picture.
 This text reviews several aspects of the pulmonary neuroendocrine system:
 phylogeny, the anine and peptide content of its epithelial cells, ontogeny
 and influence on lung development, the influence of hypoxia and nonhypoxic
 stimuli, immunomodulatory function, innervation and pathology. Among the
 discoveries of the past decade, three stand out prominently because of
 their great significance: additional proof that the neural component of
 the pulmonary neuroendocrine system is sensory, sound experimental
 evidence that PNEC stimulate airway epithelial cell differentiation and
 the discovery of a specific membrane oxygen receptor in the PNEC.

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 FULL TEXT
 Corporation on STN
 ACCESSION NUMBER: 1996:623821 SCISEARCH
 THE GENUINE ARTICLE: 1091Q
 TITLE: Phosphodiesterase inhibitors suppress alpha(2)-
 adrenoreceptor-mediated 5-hydroxytryptamine release from
 airway mucosa
 AUTHOR: Freitag A; Wessler I; Racke K (Reprint)
 CORPORATE SOURCE: Univ Bonn, Inst Pharmacol & Toxicol, Reuterstr 2B, D-53113
 Bonn, Germany (Reprint); Univ Bonn, Inst Pharmacol &
 Toxicol, D-53113 Bonn, Germany; Univ Mainz, Dept
 Pharmacol, D-55101 Mainz, Germany
 COUNTRY OF AUTHOR: Germany
 SOURCE: EUROPEAN JOURNAL OF PHARMACOLOGY, (31 JUL 1998) Vol. 354,
 No. 1, pp. 67-71.
 PUBLISHER: ELSEVIER SCIENCE BV, PO BOX 211, 1000 AE AMSTERDAM,
 NETHERLANDS.
 DOCUMENT TYPE: Article; Journal
 LANGUAGE: English
 REFERENCE COUNT: 21
 ENTRY DATE: Entered STN: 1998
 Last Updated on STN: 1998

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.
 AB The outflow of 5-hydroxytryptamine (5-HT) from isolated tracheae of
 newborn rabbits was determined by high pressure liquid chromatography with
 electrochemical detection. This 5-HT outflow reflects release from
 neuroendocrine epithelial cells of the airway mucosa, as previously shown.
 Phenylephrine, via alpha(2B)-adrenoreceptors, caused a transient increase in

5-HT outflow, maximally by about 250%, an effect mediated by liberation of
 intracellular Ca2+ previously shown. The non-selective
 phosphodiesterase inhibitor 2-isobutyl-1-methylxanthine (IBMX)
 concentration-dependently inhibited phenylephrine-induced 5-HT release
 (completely at 100 mu M, IC50: 1.3 mu M). Likewise, benzazodan (inhibitor of
 phosphodiesterase 3) also almost completely inhibited phenylephrine-
 induced 5-HT release with IC50 values of 1.7 and 4.2 mu M, respectively.
 Rolipram (inhibitor of phosphodiesterase 4), in a concentration of 10 mu
 M, which exceeds more than 10-fold the reported IC50 for phosphodiesterase
 4, did not significantly affect phenylephrine-induced 5-HT release. 5-HT
 release induced by depolarizing concentrations of K+ (45 mM), which
 largely depends on extracellular Ca2+, was not affected by IBMX. In
 conclusion, phosphodiesterases, with characteristics of phosphodiesterase
 3, appear to play an important role in the control of cyclic nucleotide
 mediated inhibition of 5-HT release from neuroendocrine epithelial cells.
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 FULL TEXT
 Corporation on STN
 ACCESSION NUMBER: 1996:13500 SCISEARCH
 THE GENUINE ARTICLE: Y1923
 TITLE: Nitric oxide, via activation of guanylyl cyclase,
 suppresses alpha(2)-adrenoreceptor-mediated
 5-hydroxytryptamine release from neuroendocrine epithelial
 cells of rabbit tracheae
 AUTHOR: Freitag A; Wessler I; Racke K (Reprint)
 CORPORATE SOURCE: Univ Bonn, Inst Pharmacol & Toxicol, Reuterstr 2B, D-53113
 Bonn, Germany (Reprint); Univ Bonn, Inst Pharmacol &
 Toxicol, D-53113 Bonn, Germany; Univ Mainz, Dept
 Pharmacol, D-55101 Mainz, Germany
 COUNTRY OF AUTHOR: Germany
 SOURCE: NAUNYN-SCHMIEDEBERG ARCHIVES OF PHARMACOLOGY, (DEC 1997)
 Vol. 356, No. 6, pp. 856-859.
 PUBLISHER: SPRINGER-VERLAG, 175 FIFTH AVE, NEW YORK, NY 10010 USA.
 DOCUMENT TYPE: Article; Journal
 LANGUAGE: English
 REFERENCE COUNT: 29
 ENTRY DATE: Entered STN: 1998
 Last Updated on STN: 1998

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.
 AB Isolated tracheae of newborn rabbits were incubated in vitro and the
 outflow of 5-hydroxytryptamine (5-HT) was determined by HPLC with
 electrochemical detection. Evidence has previously been provided that this
 5-HT outflow derives from neuroendocrine epithelial (NEE) cells of the
 airway mucosa. Phenylephrine, at a maximally effective concentration of
 10 mu M, caused a transient increase in 5-HT outflow by about 250%. An
 effect mediated by alpha(2B)-adrenoreceptors, as previously shown. The
 phenylephrine-induced 5-HT release remained unchanged in calcium-free
 medium, but was reduced by 75% when the tracheae were incubated in
 calcium-free medium which contained 0.5 mM EDTA, a treatment known to
 lower also intracellular calcium. The NO donor SNAP (S-nitroso-N-acetyl-
 penicillamine, 10 mu M) almost completely inhibited phenylephrine-
 induced 5-HT release. The inhibitory effect of SNAP was prevented by ODO,
 (1H-[1, 2, 4]oxadiazolo[4, 3-a]quinoxalin-1-one), an inhibitor of soluble
 guanylyl cyclase. In contrast, 5-HT release induced by depolarizing
 concentrations of potassium (45 mM), which was reduced by 96% in
 calcium-free medium, was not affected by SNAP. In conclusion, NO, via
 activation of soluble guanylyl cyclase, inhibits 5-HT release from NEE
 cells in a stimulus-dependent manner, alpha(2)-Adrenoreceptor-mediated 5-HT
 release, which appears to be triggered by liberation of calcium from
 intracellular stores, is suppressed by NO, whereas high potassium-evoked
 5-HT release which is triggered by calcium influx through voltage
 regulated channels, is not affected.

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 L11 727 TSUTSUMI Y7/AU
 => s l11 and calcitonin

SOURCE: ACTA PATHOLOGICA JAPONICA, (APR 1992) Vol. 42, No. 4, pp. 266-294
 PUBLISHER: BLACKWELL SCIENCE, 54 UNIVERSITY ST, P O BOX 378, CARLTON VICTORIA 3053, AUSTRALIA.
 DOCUMENT TYPE: Article; Journal
 FILE SEGMENT: LIFE
 LANGUAGE: English
 REFERENCE COUNT: 30
 ENTRY DATE: Entered STN: 1994
 Last Updated on STN: 1994

AB 43-year-old woman is reported. The 47 x 45 x 33 mm tumor, located at the periphery of the S8 segment of the resected left lower lobe, revealed by immunohistochemical studies and focal venous involvement predominantly showed immunoreactivity of cytokeratin, neuron-specific enolase, cystatin C, chromogranin A, calcitonin and neurokinin B (NPY). Fewer cells were immunoreactive for calcitonin and neurokinin B (NPY). The alpha-subunit of human chorionic gonadotropin, gastrin-releasing peptide, serotonin, methionine-enkephalin and gastrin. Immunoreactive CGRP or NPY were co-localized in calcitonin-positive cells. The amyloid substance was positively labeled only for CGRP. Immunostaining for amylin, a polypeptide showing a 50% homology with CGRP, was negative. The specificity of immunostaining for calcitonin, CGRP and amylin was confirmed by immunoblotting tests using synthetic human antigens. Immunoelectron microscopic studies disclosed peptide localization in neurosecretory-type granules and CGRP immunoreactivity in extracellular amyloid fibrils. This is the first report describing CGRP as a component of amyloid of endocrine origin.

L12 ANSWER 3 OF 6 SCISEARCH COPYRIGHT (c) 2007 The Thomson Full Text
 Corporation on STN
 ACCESSION NUMBER: 1991:610361 SCISEARCH
 THE GENUINE ARTICLE: G818
 TITLE: NEUROENDOCRINE CARCINOMA OF THE URINARY-BLADDER - CASE-REPORT AND REVIEW OF THE LITERATURE
 AUTHOR: LERTIPRASERTSUK N (Reprint); TSUTSUMI Y
 CORPORATE SOURCE: TOKAI UNIV, SCH MED, DEPT PATHOL, BOHSEIDAI, ISEHARA 25911, JAPAN
 COUNTRY OF AUTHOR: JAPANESE JOURNAL OF CLINICAL ONCOLOGY, (JUN 1991) Vol. 21, No. 3, pp. 201-210.
 SOURCE: FOUNDATION FOR PROMOTION OF CANCER RESEARCH, NATL CANCER CENTER HOSPITAL 1-1 TSUKIJI 5-CHOME CHUO-KU, TOKYO 104, JAPAN.

PUBLISHER: Article; Journal
 DOCUMENT TYPE: English
 FILE SEGMENT: No References Keyed
 LANGUAGE: Entered STN: 1994
 REFERENCE COUNT: Last Updated on STN: 1994

AB A 63-year-old Japanese man complained of hematuria and pollakiuria for several months. Computed tomography and cystography disclosed an infiltrative tumor mass in the irregularly thickened apical and posterior walls of the urinary bladder. Narrowing of the vesical lumen and posterior extension of the tumor into the pelvic cavity were also noted. After palliative ureterocutaneousostomy, 60 Gy irradiation was given locally. The patient died of cachexia seven months later. Autopsy revealed neuroendocrine carcinoma of the urinary bladder with extensive invasions and metastases to the pelvic and peritoneal cavities, liver, lungs, vertebrae, left kidney and retroperitoneal lymph nodes. Histologically, atypical tumor cords with eosinophilic cytoplasm formed solid nests and anastomosing cords with pseudoglandular structures. No other histologic tumor components were included. An intact urachal remnant was found at the vesical apex while features of metaplastic cystitis were absent. In addition to positive carcinoembryonic antigen and cytokeratin, the argyrophilic cancer cells were immunoreactive for neuron-specific enolase.

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 Full Text
 L12 ANSWER 1 OF 6 SCISEARCH COPYRIGHT (c) 2007 The Thomson
 Corporation on STN
 ACCESSION NUMBER: 2001:571761 SCISEARCH
 THE GENUINE ARTICLE: 449UF

TITLE: Osteoclast-like cells in an in vitro model of bone destruction by rheumatoid synovium
 AUTHOR: Suzuki Y (Reprint); Tsutsumi Y; Nakagawa M; Suzuki H; Matsushita K; Beppu M; Aoki H; Ichikawa Y; Mizushima Y
 CORPORATE SOURCE: St Marianna Univ, Sch Med, Dept Rheumatol, Miyamae Ku, 2-16-1 Sugao, Kawasaki, Kanagawa 2168512, Japan (Reprint); St Marianna Univ, Sch Med, Dept Rheumatol, Miyamae Ku, Kawasaki, Kanagawa 2168512, Japan; St Marianna Univ, Sch Med, Inst Med Sci, Miyamae Ku, Kawasaki, Kanagawa 2168512, Japan; St Marianna Univ, Sch Med, Dept Orthoped Surg, Miyamae Ku, Kawasaki, Kanagawa 2168512, Japan; Tokai Univ, Sch Med, Dept Pathol, Isehara, Kanagawa 25911, Japan
 COUNTRY OF AUTHOR: JAPANESE JOURNAL OF CLINICAL ONCOLOGY, (JUN 2001) Vol. 40, No. 6, pp. 673-682.
 SOURCE: ISSN: 1462-0324.
 PUBLISHER: OXFORD UNIV PRESS, GREAT CLARENDON ST, OXFORD OX2 6DP, ENGLAND.
 DOCUMENT TYPE: Article; Journal
 LANGUAGE: English
 REFERENCE COUNT: 36
 ENTRY DATE: Entered STN: 27 Jul 2001
 Last Updated on STN: 27 Jul 2001

AB Objective. Osteoclasts may be involved in the process of rheumatoid bone destruction. To test this hypothesis, we developed an in vitro model of bone destruction by osteoclast-like cells derived from cultured rheumatoid synovial tissue without using any inducers.
 Methods. Synovial tissues were obtained from rheumatoid arthritis and osteoarthritis patients and tissue pieces of about 2 mm(3) that contacted synovial lining were cultured. Multinucleated cells derived from cultured synovial tissues were studied cytochemically and morphologically for osteoclast-specific markers.
 Results. Fibroblast-like and macrophage-like cells from the tissue pieces proliferated in the coexistence of lymphocytes. After 14 days of culture, multinucleated cells with tartrate-resistant acid phosphatase activity appeared. These cells expressed vacuolar H⁺-ATPase. The vitronectin receptor and cathepsin K. Although binding of 125I-labelled salmon calcitonin was very low, the cells contained ringed structures of proliferation and macrophage-like cells and formation of multinucleated cells continued during 6 months of culture in the presence of fibroblast-like cells. The bone-resorbing activity of multinucleated cells derived from rheumatoid synovial tissue was much higher than that of cells from osteoarthritis synovial tissue, and was related to the disease activity of rheumatoid arthritis.
 Conclusion. Our culture system reproduced in vitro the process of bone destruction by rheumatoid synovium, including the proliferation and fusion of precursor cells, polarization, activation and bone tissue resorption. This system may provide a tool for understanding the mechanisms of bone destruction in rheumatoid arthritis and for the development of new therapies to prevent bone destruction.

L12 ANSWER 2 OF 6 SCISEARCH COPYRIGHT (c) 2007 The Thomson Full Text
 Corporation on STN
 ACCESSION NUMBER: 1992:314509 SCISEARCH
 THE GENUINE ARTICLE: HU235
 TITLE: ATYPICAL CARCINOID-TUMOR OF THE LUNG WITH AMYLOID STROMA
 AUTHOR: ABE Y (Reprint); UTSUNOMIYA H; TSUTSUMI Y
 CORPORATE SOURCE: TOKAI UNIV, SCH MED, DEPT PATHOL, ISEHARA, KANAGAWA 25911, JAPAN (Reprint)
 COUNTRY OF AUTHOR: JAPAN

AB A 63-year-old Japanese man complained of hematuria and pollakiuria for several months. Computed tomography and cystography disclosed an infiltrative tumor mass in the irregularly thickened apical and posterior walls of the urinary bladder. Narrowing of the vesical lumen and posterior extension of the tumor into the pelvic cavity were also noted. After palliative ureterocutaneousostomy, 60 Gy irradiation was given locally. The patient died of cachexia seven months later. Autopsy revealed neuroendocrine carcinoma of the urinary bladder with extensive invasions and metastases to the pelvic and peritoneal cavities, liver, lungs, vertebrae, left kidney and retroperitoneal lymph nodes. Histologically, atypical tumor cords with eosinophilic cytoplasm formed solid nests and anastomosing cords with pseudoglandular structures. No other histologic tumor components were included. An intact urachal remnant was found at the vesical apex while features of metaplastic cystitis were absent. In addition to positive carcinoembryonic antigen and cytokeratin, the argyrophilic cancer cells were immunoreactive for neuron-specific enolase.

ENTRY DATE: Entered STN: 9 Jun 2003
 Last Updated on STN: 9 Jun 2003
 ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.
 AB In the present review we will summarize the current knowledge about the cells comprising the Diffuse Endocrine System (DES) in mammalian organs. We will describe the morphological, histochemical and functional traits of these cells in three major systems: gastrointestinal, respiratory and prostatic. We will also focus on some aspects of their ontogeny and differentiation, as well as to their relevance in carcinogenesis, especially in neuroendocrine tumors. The first chapter describes the characteristics of DES cells and some of their specific biological and gastrointestinal traits. The second chapter deals with DES in the gastrointestinal organs, with special reference to the new data on the differentiation mechanisms that leads to the appearance of endocrine cells from an undifferentiated stem cell. The third chapter is devoted to DES of the respiratory system and some aspects of its biological role, both during development and adulthood. Neuroendocrine hyperplasia and neuroendocrine lung tumors are also addressed. Finally, the last chapter deals with the prostatic DES, discussing its probable functional role and its relevance in hormone-resistant prostatic carcinomas.

AUTHOR: Kurabuchi S; Tanaka S (Reprint)
 CORPORATE SOURCE: Univ Shizuoka, Dept Biol, Fac Sci, Ohya 836, Shizuoka 4228529, Japan (Reprint); Univ Shizuoka, Dept Biol, Fac Sci, Shizuoka 4228529, Japan; Nippon Dent Univ Tokyo, Dept Histol, Sch Dent, Tokyo, Japan
 COUNTRY OF AUTHOR: Japan
 SOURCE: JOURNAL OF HISTOCHEMISTRY & CYTOCHEMISTRY, (JUL 2002) Vol. 50, No. 7, pp. 903-909.
 ISSN: 0022-1554.
 PUBLISHER: HISTOCHEMICAL SOC INC, UNIV WASHINGTON, DEPT BIOSTRUCTURE, BOX 357420, SEATTLE, WA 98195 USA.
 DOCUMENT TYPE: Article; Journal
 LANGUAGE: English
 REFERENCE COUNT: 36
 ENTRY DATE: Entered STN: 19 Jul 2002
 Last Updated on STN: 19 Jul 2002
 ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.

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AUTHOR: Kurabuchi S; Tanaka S (Reprint)
 CORPORATE SOURCE: Univ Shizuoka, Dept Biol, Fac Sci, Ohya 836, Shizuoka 4228529, Japan (Reprint); Univ Shizuoka, Dept Biol, Fac Sci, Shizuoka 4228529, Japan; Nippon Dent Univ Tokyo, Dept Histol, Sch Dent, Tokyo, Japan
 COUNTRY OF AUTHOR: Japan
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 PUBLISHER: HISTOCHEMICAL SOC INC, UNIV WASHINGTON, DEPT BIOSTRUCTURE, BOX 357420, SEATTLE, WA 98195 USA.
 DOCUMENT TYPE: Article; Journal
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 CORPORATE SOURCE: Univ Shizuoka, Dept Biol, Fac Sci, Ohya 836, Shizuoka 4228529, Japan (Reprint); Univ Shizuoka, Dept Biol, Fac Sci, Shizuoka 4228529, Japan; Nippon Dent Univ Tokyo, Dept Histol, Sch Dent, Tokyo, Japan
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 DOCUMENT TYPE: Article; Journal
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AUTHOR: Kurabuchi S; Tanaka S (Reprint)
 CORPORATE SOURCE: Univ Shizuoka, Dept Biol, Fac Sci, Ohya 836, Shizuoka 4228529, Japan (Reprint); Univ Shizuoka, Dept Biol, Fac Sci, Shizuoka 4228529, Japan; Nippon Dent Univ Tokyo, Dept Histol, Sch Dent, Tokyo, Japan
 COUNTRY OF AUTHOR: Japan
 SOURCE: JOURNAL OF HISTOCHEMISTRY & CYTOCHEMISTRY, (JUL 2002) Vol. 50, No. 7, pp. 903-909.
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 PUBLISHER: HISTOCHEMICAL SOC INC, UNIV WASHINGTON, DEPT BIOSTRUCTURE, BOX 357420, SEATTLE, WA 98195 USA.
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 ENTRY DATE: Entered STN: 19 Jul 2002
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AUTHOR: Kurabuchi S; Tanaka S (Reprint)
 CORPORATE SOURCE: Univ Shizuoka, Dept Biol, Fac Sci, Ohya 836, Shizuoka 4228529, Japan (Reprint); Univ Shizuoka, Dept Biol, Fac Sci, Shizuoka 4228529, Japan; Nippon Dent Univ Tokyo, Dept Histol, Sch Dent, Tokyo, Japan
 COUNTRY OF AUTHOR: Japan
 SOURCE: JOURNAL OF HISTOCHEMISTRY & CYTOCHEMISTRY, (JUL 2002) Vol. 50, No. 7, pp. 903-909.
 ISSN: 0022-1554.
 PUBLISHER: HISTOCHEMICAL SOC INC, UNIV WASHINGTON, DEPT BIOSTRUCTURE, BOX 357420, SEATTLE, WA 98195 USA.
 DOCUMENT TYPE: Article; Journal
 LANGUAGE: English
 REFERENCE COUNT: 36
 ENTRY DATE: Entered STN: 19 Jul 2002
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 ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.

ENTRY DATE: Entered STN: 19 Jul 2002
 Last Updated on STN: 19 Jul 2002
 ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.
 AB We examined the immunocytochemical localization of the proenzyme convertases PC1 and PC2 in the thyroid gland and renal pheochromocytoma of the adult mouse using the indirect enzyme- and immunogold-labeled antibody methods for light and electron microscopy, respectively. In the thyroid gland, PC1- and/or PC2-immunoreactive cells were cuboidal, scattered in the follicular epithelium and in the interfollicular spaces. When serial sections were immunostained with anti-calcitonin, anti-PC1, anti-calcitonin-gene-related-peptide (CGRP), and anti-PC2 sera, respectively, localization of both PC1 and PC2 was restricted to the calcitonin/CGRP-producing parafollicular cells. In the respiratory tract, only PC1 immunoreactivity was observed in the basal granulated neuroendocrine cells, which were scattered in the tracheal epithelium. Consecutive sections immunostained with anti-PC1 and anti-CGRP sera showed that a subpopulation of these PC1-immunoreactive cells contained CGRP. Double immunogold electron microscopy of the thyroid parafollicular cells revealed that calcitonin- and/or CGRP-immunopositive secretory granules were also labeled with both PC1 and PC2. These findings suggest that procalcitonin is proteolytically cleaved by PC2 alone or by PC2 together with PC1, and that the proCGRP is cleaved by PC1.

AUTHOR: Kurabuchi S; Tanaka S (Reprint)
 CORPORATE SOURCE: Univ Shizuoka, Dept Biol, Fac Sci, Ohya 836, Shizuoka 4228529, Japan (Reprint); Univ Shizuoka, Dept Biol, Fac Sci, Shizuoka 4228529, Japan; Nippon Dent Univ Tokyo, Dept Histol, Sch Dent, Tokyo, Japan
 COUNTRY OF AUTHOR: Japan
 SOURCE: JOURNAL OF HISTOCHEMISTRY & CYTOCHEMISTRY, (JUL 2002) Vol. 50, No. 7, pp. 903-909.
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 PUBLISHER: HISTOCHEMICAL SOC INC, UNIV WASHINGTON, DEPT BIOSTRUCTURE, BOX 357420, SEATTLE, WA 98195 USA.
 DOCUMENT TYPE: Article; Journal
 LANGUAGE: English
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 ENTRY DATE: Entered STN: 19 Jul 2002
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AUTHOR: Kurabuchi S; Tanaka S (Reprint)
 CORPORATE SOURCE: Univ Shizuoka, Dept Biol, Fac Sci, Ohya 836, Shizuoka 4228529, Japan (Reprint); Univ Shizuoka, Dept Biol, Fac Sci, Shizuoka 4228529, Japan; Nippon Dent Univ Tokyo, Dept Histol, Sch Dent, Tokyo, Japan
 COUNTRY OF AUTHOR: Japan
 SOURCE: JOURNAL OF HISTOCHEMISTRY & CYTOCHEMISTRY, (JUL 2002) Vol. 50, No. 7, pp. 903-909.
 ISSN: 0022-1554.
 PUBLISHER: HISTOCHEMICAL SOC INC, UNIV WASHINGTON, DEPT BIOSTRUCTURE, BOX 357420, SEATTLE, WA 98195 USA.
 DOCUMENT TYPE: Article; Journal
 LANGUAGE: English
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 ENTRY DATE: Entered STN: 19 Jul 2002
 Last Updated on STN: 19 Jul 2002
 ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.

hypertension. Pulmonary neuroendocrine cells (PNEC) produce calcitonin gene-related peptide (CGRP), a potent vasodilator. We previously reported altered distribution of CGRP-positive PNEC in full-term rats with CDH, that may lead to an imbalance in vasoactive mediators. In the present study we examined the expression of CGRP-positive PNEC during lung development in rats with CDH induced by 2,4-dichlorophenyl-p-nitrophenyl (2,4-DCPF) and the lungs were quantitated through image analysis. On Day 16, CGRP-immunoreactive staining was negative. On Day 22, CGRP-immunoreactive staining was positive in all controls (not exposed to Nitrofen), whereas in CDH pups CGRP-positive cells were present in only four of six cases. On Day 20, CGRP immunoreactivity was similar in CDH pups. Nitrofen-exposed pups without CDH and controls. On Day 22 (term), significantly more CGRP-positive cells (i.e., number of positive cells per surface area [mm²] or lung volume [mm³]) were found in ipsilateral lungs of CDH pups than in controls ($P < 0.05$). The difference was even more striking in contralateral lungs of CDH pups ($P < 0.001$), ruling out nonspecific effects of Nitrofen. In CDH lungs, the proportion of immunostained epithelium and the size of the neuroendocrine cell clusters (neuroepithelial bodies [NEB]) were not significantly different from those of controls. On Day 22, supraoptimal dilution immunocytochemistry yielded similar results in CDH pups and controls. We conclude that in CDH, CGRP expression in PNEC and NEB is delayed during early stages of lung development. Because CGRP also exhibits growth factor-like properties for endothelium and epithelial cells, the lack of this factor during a crucial developmental stage (canalicular period) may be causally related to lung hypoplasia.

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Full Text
 Corporation on STN
 Accession Number: 1995:199991 SCISEARCH
 The Genuine Article: QM211
 Title: PULMONARY NEUROENDOCRINE CELLS IN NEONATAL RATS WITH CONGENITAL DIAPHRAGMATIC HERNIA
 Author: IJSELSIJN H (Reprint); PERRIN D G; DEJONGSTE J C; CUTZ E; TIBBOEL D
 Corporate Source: SOPHIA CHILDRENS UNIV HOSP, DEPT PEDIAT SURG, 3015 GJ ROTTERDAM, NETHERLANDS; ERASMUS UNIV ROTTERDAM, DEPT PEDIAT SURG, 3000 D ROTTERDAM, NETHERLANDS; ERASMUS UNIV ROTTERDAM, DEPT PEDIAT, DIV RESP MED, ROTTERDAM, NETHERLANDS; HOSP SICK CHILDREN, RES INST, MRC, LUNG DEV GRP, TORONTO, ON M5G 1X8, CANADA; HOSP SICK CHILDREN, DEPT PATHOL, MRC, LUNG DEV GRP, TORONTO, ON M5G 1X8, CANADA
 Country of Author: NETHERLANDS; CANADA
 Source: JOURNAL OF PEDIATRIC SURGERY, (MAR 1995) Vol. 30, No. 3, pp 413-415
 ISSN: 0022-3468
 Publisher: W B SAUNDERS
 Document Type: Article; Journal
 File Segment: English
 Language: English
 Reference Count: 20
 Entry Date: Entered STN: 1995

*ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.
 AB Lung hypoplasia and persistent pulmonary hypertension are the principal causes of high mortality and morbidity in infants with congenital diaphragmatic hernia (CDH). Amine and peptide producing pulmonary neuroendocrine cells (PNEC), widely distributed throughout the airway mucosa, are thought to play an important role in both pulmonary development and regulation of pulmonary vascular tone. Furthermore, recent studies show increased levels of calcitonin gene-related peptide (CGRP), a pulmonary vasodilator produced by PNEC, during chronic hypoxia. The article reports data on morphometric analysis of CGRP immunoreactive PNEC clusters (neuroepithelial bodies, NEB) in a rat model of CDH. CDH was induced in neonatal Sprague Dawley rats by oral administration of 2,4-dichlorophenyl-p-nitrophenyl ether (Nitrofen; Rohm Haas, Philadelphia, PA) to the mother at 10 days of gestation. Sections of lungs from term neonatal rats with and without CDH and controls were immunostained for

CGRP (marker of NEB) with specific antibody against rat CGRP. NEB size and number of NEB/area of lung were assessed using a semiautomatic image analysis system. In lungs of neonatal rats with CDH, the number of NEB per surface area of lung parenchyma was significantly increased compared with the age-matched controls. Although the mean size of NEB was larger in CDH lungs, the size of NEB was not significantly different from controls. PNEC in CDH lungs were not significantly increased in this study, but altered NEB function, including imbalance in vasoactive mediators, requires further studies, especially in the human being. Copyright (c) 1995 by W.B. Saunders Company

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Full Text
 Corporation on STN
 Accession Number: 1994:424640 SCISEARCH
 The Genuine Article: NV118
 Title: SMALL-CELL LUNG-CARCINOMA CELL-LINES EXPRESS MESSENGER-RNA FOR CALCITONIN AND ALPHA-CALCITONIN AND BETA-CALCITONIN GENE-RELATED PEPTIDES
 Author: KELLEY M J (Reprint); SNIDER R H; BECKER K L; JOHNSON B E
 Corporate Source: NCI, USN, MED ONCOL BRANCH, BLDG 8, ROOM 5101, BETHESDA, MD 20889 (Reprint); VET AFFAIRS MED CTR, WASHINGTON, DC 20422; GEORGE WASHINGTON UNIV, WASHINGTON, DC 20037
 Country of Author: USA
 Source: CANCER LETTERS, (15 JUN 1994) Vol. 81, No. 1, pp. 19-25. ISSN: 0304-3835
 Publisher: ELSEVIER SCI IRELAND LTD, CUSTOMER RELATIONS MANAGER, BAY 15, SHANNON INDUSTRIAL ESTATE CO, CLARE, IRELAND.
 Document Type: Article; Journal
 File Segment: English
 Language: English
 Reference Count: 23
 Entry Date: Entered STN: 1994

*ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.
 AB Calcitonin (CT) and calcitonin gene related peptide (CGRP) are derived from preprohormones encoded by three mRNAs (CT, alpha-CGRP and beta-CGRP) from two genes (CALC1 and CALC2) on chromosome 11. Among 16 small cell lung cancer cell lines examined by RNase protection assay, 9 (56%) had detectable CT mRNA, 8 (50%) had alpha-CGRP mRNA, and 13 (81%) had beta-CGRP mRNA. At least one CALC1 transcript (CT or alpha-CGRP) was found in 11 (69%) cell lines with three having only CT mRNA, two having only alpha-CGRP mRNA, and six having both. beta-CGRP mRNA was detected in all of these 11 cell lines expressing a CALC1 transcript. Immunoreactive CT was detected by radioimmunoassay in eight of nine SCJC cell lines expressing CT mRNA, and immunoreactive CGRP was detected in 12 of 13 cell lines expressing a CGRP mRNA. The variety of expression of these three peptides in different cell lines of the same cell type should provide a useful system for further study of the control of expression of these peptides.

L15 ANSWER 7 OF 14 SCISEARCH COPYRIGHT (c) 2007 The Thomson

Full Text
 Corporation on STN
 Accession Number: 1994:378457 SCISEARCH
 The Genuine Article: NP717
 Title: CALCITONIN ELEVATION IN SMALL-CELL LUNG-CANCER WITHOUT ECTOPIC PRODUCTION
 Author: KELLEY M J (Reprint); BECKER K L; RUSHIN J M; VENZON D; PHELPS R; INDE D C; BLISS D P; MELBY K; SNIDER R H; JOHNSON B E
 Corporate Source: NATL NAVAL MED CTR, NCI, NAVY MED ONCOL BRANCH, BLDG 8, ROOM 5101, BETHESDA, MD 20889 USA (Reprint); NCI, BIOTAT & DATA MANAGEMENT SECT, BETHESDA, MD USA; VET ADM MED CTR, DIV ENDOCRINOL, WASHINGTON, DC 20422 USA; GEORGE WASHINGTON UNIV, WASHINGTON, DC USA; ST JOSEPHS HOSP, ATLANTA, GA USA; NATL NAVAL MED CTR, LAB DEPT, BETHESDA, MD USA
 Country of Author: USA
 Source: AMERICAN JOURNAL OF RESPIRATORY AND CRITICAL CARE MEDICINE (JAN 1994) Vol. 149, No. 1, pp. 183-190. ISSN: 1073-449X.

PUBLISHER: AMER THORACIC SOC, 1740 BROADWAY, NEW YORK, NY 10019-4374
DOCUMENT TYPE: Article; Journal
LANGUAGE: English
REFERENCE COUNT: 39
ENTRY DATE: Entered STN: 1994
 Last Updated on STN: 31 AUG 2006

AB To determine the relative contribution of ectopic calcitonin (CT) production versus nonectopic secretion of CT in patients with small cell lung cancer (SCLC) serum and urine immunoreactive CT (iCT) levels of 86 different subjects were measured by radioimmunoassay (RIA) using two polyclonal antisera (Ab3b and Ab4). The subjects included 49 previously untreated patients with SCLC, 17 smokers, and 20 nonsmokers. Serum and urine iCT values were highest in the patients with SCLC, intermediate in the smokers, and lowest in the nonsmokers ($p < 0.0003$). Sixteen of the 49 patients with SCLC had tumor cell lines available for determination of CT mRNA expression by RNase protection assay (RPA) and iCT production by RIA. CT mRNA was detected in nine of 16 subjects and iCT in eight of 16. The tumor cell lines of seven patients had undetectable CT by both RPA and RIA, and of these, five had elevated urine or serum iCT values compared with those of nonsmokers, and two had levels above all values in the smoker group. Immunohistochemical staining of formalin-fixed, paraffin-embedded tumor samples detected iCT in two of four tumors from patients whose tumor cell lines had CT mRNA by RPA and iCT by RIA, but in none of six whose tumor cell lines had undetectable CT mRNA. Thus, increased iCT values in some patients with SCLC are likely due to sources other than CT production by tumor cells.

L15 ANSWER 8 OF 14 SCISEARCH COPYRIGHT (c) 2007 The Thomson
Full Text
Accession Number: STN Corporation on 1994:298011 SCISEARCH
The Genuine Article: NK671
Title: DETECTION OF CALCITONIN-GENE EXPRESSION IN HUMAN INFANT AND MONKEY CAROTID-BODY CHIEF CELLS BY IN-SITU HYBRIDIZATION
Author: WANG Y Y (Reprint); CUTZ E; PERRIN D G
Corporate Source: HOSP SICK CHILDREN, DEPT PATHOL, TORONTO M5G 1X8, ON, CANADA; HOSP SICK CHILDREN, RES INST, TORONTO M5G 1X8, ON, CANADA; UNIV TORONTO, TORONTO, ON, CANADA
Country of Author: CANADA
Source: CELL AND TISSUE RESEARCH, (MAY 1994) Vol. 276, No. 2, pp. 399-402.
 ISSN: 0302-766X
 SPRINGER, 233 SPRING STREET, NEW YORK, NY 10013 USA.
Publisher: Article; Journal
Document Type: English
Reference Count: 26
Entry Date: Entered STN: 1994
 Last Updated on STN: 1994

AB Calcitonin mRNA was detected in human and monkey carotid bodies by in situ hybridization histochemistry using a S-35-labeled oligonucleotide probe for human calcitonin. In both human and monkey carotid body, moderate to high hybridization signal for calcitonin mRNA was observed in all cases. The hybridization signal in the formalin-fixed, paraffin-embedded samples was comparable to that obtained from frozen paraformaldehyde-fixed tissue. Our observations extend the finding of calcitonin-like immunoreactivity in the carotid body chief cells and indicate that calcitonin is produced in the carotid body, probably in the chief cells.

L15 ANSWER 9 OF 14 SCISEARCH COPYRIGHT (c) 2007 The Thomson
Full Text
Accession Number: STN Corporation on 1993:420598 SCISEARCH
The Genuine Article: LK721
Title: CALCITONIN-GENE-RELATED PEPTIDE IN SMALL-CELL LUNG CARCINOMAS
Author: SCHIFTER S (Reprint); JOHANNSEN L; BUNKER C; BRICKELL P; BORK E; LINDEBERG H; FABER J

CORPORATE SOURCE: GLOSTRUP UNIV HOSP, DEPT CLIN PHYSIOL & NUCL MED, DK-2600 GLOSTRUP, DENMARK (Reprint); RIGSHOSP, DEPT CLIN PHYSIOL & NUCL MED, DK-2100 COPENHAGEN, DENMARK; OENSEN HOSP, DEPT CLIN PHYSIOL & NUCL MED, DK-5000 OENSEN, DENMARK; UNIV COLLEGE HOSP, DEPT NEUROL, DK-5000 OENSEN, DENMARK; UNIV COLLEGE HOSP, DEPT MED, DEPT BIOCHEM MED, NO-0407 OSLO, NORWAY; UNIV LONDON WIP, ED, ENGLAND; BISPELBERG HOSP, DEPT INTERNAL MED C, DK-2400 COPENHAGEN, DENMARK; HERLEV UNIV HOSP, DEPT ENDOCRINOL F, DK-2730 HERLEV, DENMARK; AARHUS UNIV HOSP, DEPT OTOLARYNGOL, DK-8000 AARHUS, DENMARK
Country of Author: DENMARK; ENGLAND
Source: CLINICAL ENDOCRINOLOGY, (JUL 1993) Vol. 39, No. 1, pp. 59-65
 ISSN: 0300-0664
 BLACKWELL SCIENCE LTD, OSNEY MEAD, OXFORD, OXON, ENGLAND
Publisher: Article; Journal
Document Type: English
File Segment: LIFE; CLIN
Language: English
Reference Count: 31
Entry Date: Entered STN: 1994
 Last Updated on STN: 1994

AB OBJECTIVE Calcitonin gene-related peptide (CGRP) is a regulatory peptide encoded by the calcitonin gene. CGRP is expressed in increased amounts by the cells of medullary thyroid carcinomas and has been demonstrated by immunohistochemistry to occur in neuroendocrine cells and nerve fibres of lung tissue.
 MEASUREMENTS Serum CGRP levels were measured in patients with small cell lung carcinomas before treatment (n=74) and immediately before the second course of chemotherapy (n=30). In-situ hybridization and immunohistochemistry were performed on tumour tissue and CGRP was extracted from two tumours and characterized by gel chromatography and high pressure liquid chromatography.
 RESULTS Serum CGRP levels were elevated in small cell lung carcinomas when compared with healthy controls of similar age and sex (median values 55.0 vs 36.6 pmol/l, $P < 0.001$), and 274 had levels above the upper normal range. Serum CGRP levels decreased following the initial course of chemotherapy ($P < 0.05$) but remained elevated when compared to the controls ($P < 0.001$). In-situ hybridization for CGRP mRNA was positive in three of 17 tumours and immunohistochemistry was positive in seven of 31 tumours investigated. CGRP immunoreactivity extracted from two tumours was characterized by gel chromatography and high pressure liquid chromatography. A major part of the immunoreactivity was demonstrated to represent the intact molecule.
 CONCLUSIONS We found that patients with small cell lung carcinomas had elevated concentration of serum calcitonin gene-related peptide but only 274 had values above the upper normal range. Serum CGRP is therefore of limited value as a tumour marker. Intact CGRP can be extracted from tumour tissue, and in-situ hybridization and immunohistochemistry showed elevated expression of CGRP in a few of the tumours investigated. The elevated serum CGRP levels are therefore likely to be largely of extratumoral origin.

L15 ANSWER 10 OF 14 SCISEARCH COPYRIGHT (c) 2007 The Thomson
Full Text
Accession Number: STN Corporation on 1992:314509 SCISEARCH
The Genuine Article: HU235
Title: ATYPICAL CARCINOID-TUMOR OF THE LUNG WITH AMYLOID STROMA
Author: ABE Y (Reprint); UTSUNOMIYA H; TSUTSUMI Y
Corporate Source: TOKAI UNIV, SCH MED, DEPT PATHOL, ISEHARA, KANAGAWA 25911, JAPAN (Reprint)
Country of Author: JAPAN
Source: ACTA PATHOLOGICA JAPONICA, (APR 1992) Vol. 42, No. 4, pp. 286-292.
 ISSN: 0001-6632
 BLACKWELL SCIENCE, 54 UNIVERSITY ST, P O BOX 378, CARLTON VICTORIA 3053, AUSTRALIA.
Publisher: Article; Journal
Document Type: LIFE
File Segment: English
Language: English
Reference Count: 30

ENTRY DATE:

Entered STN: 1994

Last Updated on STN: 1994

*ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.

AB Atypical carcinoid tumor of the lung with amyloid stroma seen in a 43-year-old woman is reported. The 47 x 45 x 13 mm tumor located at the periphery of the S8 segment of the resected left lower lobe revealed a well-differentiated neuroendocrine tumor. The atypical neuroendocrine tumor cells with occasional mitoses and focal venous involvement. Predominantly chromogranin A, calcitonin, and neurokinin B (NPY) immunoreactivity were shown. Chromogranin A, calcitonin, and neurokinin B (NPY) immunoreactivity were shown. Fewer cells were immunoreactive for calcitonin gene-related peptide (CGRP), the alpha-subunit of human chorionic gonadotropin, gastrin-releasing peptide, serotonin, methionine-enkephalin and gastrin. Immunoreactive CGRP or NPY were co-localized in calcitonin-positive cells. The amyloid substance was positively labeled only for CGRP. Immunostaining for amylin, a polypeptide isolated from insulin amyloid in type II diabetes mellitus or insulinoma showing a 50% homology with CGRP, was negative. The specificity of immunostaining for calcitonin, CGRP and amylin was confirmed by immunoblotting tests using synthetic human antigens. Immunoelectron microscopic studies disclosed peptide localization in neurosecretory-type granules and CGRP immunoreactivity in extracellular amyloid fibrils. This is the first report describing CGRP as a component of amyloid of endocrine origin.

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Full Text

Corporation on STN

Accession Number: 1991:193329 SCISEARCH

The Genuine Article: HJ298

TITLES: DIFFERENTIAL DIAGNOSTIC PATTERNS OF LUNG NEUROENDOCRINE TUMORS - A CLINICOPATHOLOGICAL AND IMMUNOHISTOCHEMICAL STUDY OF 122 CASES

AUTHOR: BONATO M (Reprint); CERATI M; PAGANI A; PAPOTTI M; BOSI F; RUSSOLATI G; CAPELLA C

CORPORATE SOURCE: UNIV PAVIA, PAC MED 2, DEPT HUMAN PATHOL, VIALE BORRI 57, I-27100 VARESE, ITALY; POLICLIN SAN MATTEO, IST RICOVERO & CURA CARATTERI SCI, DEPT PATHOL, PAVIA, ITALY; UNIV TURIN, DEPT BIOMED SCI & HUMAN ONCOL, I-10124 TURIN, ITALY

COUNTRY OF AUTHOR: ITALY

SOURCE: VIKINGS ARCHIV A-PATHOLOGICAL ANATOMY AND HISTOPATHOLOGY, (MAR 1992) Vol. 420, No. 3, PP. 201-211.

PUBLISHER: SPRINGER VERLAG, 175 FIFTH AVE, NEW YORK, NY 10010.

DOCUMENT TYPE: Article; Journal

FILE SEGMENT: LIFE

LANGUAGE: English

REFERENCE COUNT: 4

ENTRY DATE: Entered STN: 1994

Last Updated on STN: 1994

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AB A series of 3 tumours (T1s) and 81 typical carcinoids (TCS) 14 atypical carcinoids (ACs) (well-differentiated neuroendocrine carcinomas, WDNCs) and 24 small cell-intermediate cell carcinomas (SCC-ICCs) of the lung were studied. Histopathological features were correlated with age and peptide hormone immunoreactivity and with clinical data. All types of tumours expressed general neuroendocrine (NE) markers: Grinellus positivity and chromogranins were detected more frequently in well-differentiated (T1s, TCS) than in less well differentiated tumours (ACs (WDNCs) and SCC-ICCs) whereas neuron specific enolase (NSE) was prominent in the latter tumours. T1s and peripheral TCS were benign, often showing a paraganglioid pattern and frequently expressing gastrin-releasing peptide (GRP), which is present in the peripheral airways of normal lung. Central TCS were associated with lymph node metastases in 8.5% of the cases, frequently had a trabecular architecture, often associated with human milk fat globule 2 (HMFG2)-positive acinar and rosette-like structures, and were mainly immunostained for the alpha-subunit of human chorionic gonadotropin (alpha-hCG) and serotonin. ACs (WDNCs) were associated with intrathoracic and/or extrathoracic metastases in 57.1% of the cases with a mortality rate of 35.7%. Their histological and cytological features were intermediate between those of TCS and SCC-ICCs. ACs (WDNCs) expressed serotonin and alpha-hCG less frequently than TCS. All SCC-ICCs were surgically treated and displayed a

mortality rate of 91.6% with a mean survival of 10.2 months after operation. These tumours were characterized by high expression of HMFG2 and NSE. While the expression of both orihotopic (serotonin, NPY) and ectopic (ACTH) specific substances was very low, since a 1% of either acropit (ACTH) or NPY had a favorable outcome, while about 16% of ACs (WDNCs) or TCS had the latter seen more frequently. The designation "well-differentiated NE carcinomas" The differential diagnosis between different NE tumours of the lung is important and is mainly based on morphology. Both panendocrine and specific immunohistochemical markers are helpful in distinguishing the less aggressive, mostly benign varieties from the more malignant varieties.

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Full Text

Corporation on STN

Accession Number: 1991:335987 SCISEARCH

The Genuine Article: FC0329

TITLES: COOCCURRENCE OF IMMUNOREACTIVE CALCITONIN AND CALCITONIN GENE-RELATED PEPTIDE IN NEUROENDOCRINE CELLS OF RAT LUNGS

AUTHOR: SHIMOSEGAWA T (Reprint); SAID S I

CORPORATE SOURCE: UNIV ILLINOIS, COLL MED, DEPT MED, 1940 W TAYLOR ST, CHICAGO, IL 60612; VET AFFAIRS W SIDE MED CTR, CHICAGO, IL 60612

COUNTRY OF AUTHOR: USA

SOURCE: CELL AND TISSUE RESEARCH, (JUN 1991) Vol. 264, No. 3, PP. 555-561.

ISSN: 0302-766X.

PUBLISHER: SPRINGER VERLAG, 175 FIFTH AVE, NEW YORK, NY 10010.

DOCUMENT TYPE: Article; Journal

FILE SEGMENT: LIFE

LANGUAGE: English

REFERENCE COUNT: 51

ENTRY DATE: Entered STN: 1994

Last Updated on STN: 1994

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AB Neuroendocrine cells of the lung, occurring singly or in clusters known as neuroepithelial bodies, contain a variety of biologically active compounds, including several neuropeptides. We have investigated the localization of calcitonin and calcitonin gene-related peptide (CGRP) within single and grouped neuroendocrine cells in the respiratory epithelium of rats by an immunohistochemical double-staining technique which uses specific antisera raised in heterogeneous animal species. Calcitonin- and CGRP-immunoreactivities were nearly totally co-localized in both single neuroendocrine cells and neuroepithelial bodies. CGRP-immunoreactivity was also present in neurons in the jugular, nodose and dorsal root ganglia. The calcitonin-immunoreactivity in neuroendocrine cells, as in thyroid parafollicular (C) cells, was abolished by the administration of the calcitonin receptor antagonist, calcitonin. The CGRP-immunoreactivity in neuroendocrine cells and in neuronal cells was abolished by the administration of anti-CGRP with synthetic CGRP. Thus while the calcitonin gene is expressed exclusively or predominantly as either calcitonin or CGRP in all other tissues except thyroid C-cells, our results strongly suggest that both peptides are expressed in the rat bronchopulmonary neuroendocrine cells.

L15 ANSWER 13 OF 14 SCISEARCH COPYRIGHT (c) 2007 The Thomson

Full Text

Corporation on STN

Accession Number: 1991:305209 SCISEARCH

The Genuine Article: FM530

TITLES: NEUROENDOCRINE TUMORS OF THE LUNG WITH PROPOSED CRITERIA FOR LARGE-CELL NEUROENDOCRINE CARCINOMA - AN ULTRASTRUCTURAL, IMMUNOHISTOCHEMICAL, AND FLOW CYTOMETRIC STUDY OF 35 CASES

AUTHOR: TRAVIS W D (Reprint); LINNOILA R I; TSOKOS M G; HITCHCOCK C L; CUTLER G B; NIEMAN L; CHROUSOS G; PASS H; DOPPMAN J

CORPORATE SOURCE: NICHID, SURG BRANCH, BETHESDA, MD 20892; NICHID, DEV ENDOCRINOL BRANCH, BETHESDA, MD 20892; NIH, WARREN G MAGNUSON CLIN CTR, DEPT DIAGNOST RADIOLOG, BETHESDA, MD 20892; ARMED FORCES INST PATHOL, DEPT CELLULAR PATHOL, WASHINGTON, DC 20306; USN HOSP, PATHOL LAB, BETHESDA, MD 20814; USN HOSP, NCI, NAVY MED ONCOL BRANCH, BETHESDA, MD

20814
 COUNTRY OF AUTHOR: USA
 SOURCE: AMERICAN JOURNAL OF SURGICAL PATHOLOGY, (JUN 1991) Vol. 15, No. 6, PP. 529-553.
 PUBLISHER: LIPPINCOTT-RAVEN PUBL, 227 EAST WASHINGTON SQ, PHILADELPHIA, PA 19106.
 DOCUMENT TYPE: Article; Journal
 FILE SEGMENT: LIFE; CLIN
 LANGUAGE: English
 REFERENCE COUNT: 91
 ENTRY DATE: Entered STN: 1994
 Last Updated on STN: 1994

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.
 AB Based on our review of 35 cases and the literature, we found the spectrum of pulmonary neuroendocrine (NE) tumors to be too broad to fit into the traditional three-category classification scheme of typical carcinoid (TC), atypical carcinoid (AC), and small-cell lung carcinoma (SCLC). We found that a spectrum of high- and low-grade tumors exist between TC and SCLC and that in the past many of these tumors have been called AC. We chose to adhere to Arigoni's definition of AC, as his original criteria characterized a low-grade tumor. For the higher grade non-small-cell tumors (NSCLC), we propose a fourth category of large-cell neuroendocrine carcinoma (LCNEC), which is characterized by: (a) light microscopic NE appearance; (b) cells of large size, polygonal shape, low nuclear-cytoplasmic ratio (N:C), coarse nuclear chromatin, and frequent nucleoli; (c) high mitotic rate (> 10/10 high-power fields (HPF)) and frequent necrosis; and (d) NE features by immunohistochemistry (IHC) or electron microscopy (EM). Thus, after deciding that a pulmonary NE tumor is high grade, the major diagnostic issue is separation of LCNEC from SCLC. This distinction is based not only on cell size, but on a variety of morphologic features. We studied 20 TC, six AC, five LCNEC, and four SCLC and characterized the clinical, light microscopic, EM, IHC, and flow cytometric features of each type of tumor. We did not find any advantage to IHC, EM, or flow cytometry over light microscopy in the subclassification or prediction of prognosis; however, these methods were useful in characterizing these four types of pulmonary NE tumors and in demonstrating their NE properties. LCNEC must be distinguished from a fifth category pulmonary NE tumor: NSCLC with NE features in which NE differentiation is not evident by light microscopy and must be demonstrated by EM or IHC. Although the prognosis of LCNEC appears to be intermediate between AC and SCLC, larger numbers of patients will be needed to demonstrate significant differences in survival.

L15 ANSWER 14 OF 14 SCISEARCH COPYRIGHT (c) 2007 The Thomson
 Full Text
 Corporation on STN
 Accession Number: 1991:183590 SCISEARCH
 THE GENUINE ARTICLE: PD026
 TITLE: TUMORLETS OF THE LUNG - AN ULTRASTRUCTURAL-STUDY
 AUTHOR: TORIKATA C (Reprint)
 CORPORATE SOURCE: KEIO UNIV, SCH MED, DEPT PATHOL, 35 SHINANO MACHI, SHINJUKU KU, TOKYO 160, JAPAN (Reprint)
 COUNTRY OF AUTHOR: JAPAN
 SOURCE: ULTRASTRUCTURAL PATHOLOGY, (MAR-APR 1991) Vol. 15, No. 2, PP. 189-195.
 ISSN: 0191-3123.
 PUBLISHER: HEMISPHERE PUBL CORP, 1900 FROST ROAD, SUITE 101, BRISTOL, PA 19007-1598.
 DOCUMENT TYPE: Article; Journal
 FILE SEGMENT: LIFE
 LANGUAGE: English
 REFERENCE COUNT: 24
 ENTRY DATE: Entered STN: 1994
 Last Updated on STN: 1994

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.
 AB A tumorlet of the lung is a minute tumorlike lesion found in damaged lungs in close association with the bronchioles. Histochemical and ultrastructural studies identify proliferating cells in the tumorlets as Kulchitzky-type cells. However, the pathological significance of the tumorlets, whether they are hyperplastic or neoplastic, is still controversial. Previous ultrastructural studies on the tumorlets have

been carried out on formalin-fixed lung tissues. The case examined in this study was of typical tumorlets found in a so-called middle lobe of the lung of a 54-year-old man. Tumorlets were located within the bronchial mucosa, extended directly by a basal lamina and by the broncholar noncondensing epithelial cells. There were no signs of invasion into the surrounding connective tissues or into lymphatic spaces. Between the covering broncholar epithelial cells and the subjacent proliferating Kulchitzky cells, specific sites of cell-to-cell attachment were noted. This finding, in addition to the previously reported clinicopathological characteristics, indicates that the proliferating Kulchitzky-type cells in the tumorlets might be non-neoplastic and that tumorlets are due to hyperplasia of pure Kulchitzky-type cells, thus resembling neuroepithelial bodies of the lung.

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 L2 1797 S GRAY M7/AU
 L3 49 S L1 AND L2
 L4 11201 S JOHNSON M7/AU
 L5 5 S L4 AND L3
 L6 SEL L5 4 CIT
 21 S E1
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 L12 727 S TSUTSUMI Y7/AU
 L13 6 S L11 AND CALCITONIN
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 L17 ANSWER 1 OF 11 SCISEARCH COPYRIGHT (c) 2007 The Thomson
 Full Text
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 Accession Number: 2007:71970 SCISEARCH

THE GENUINE ARTICLE: 121KV

TITLE: Pulmonary endocrine cells in hypoplastic lungs due to

AUTHOR: fetal urinary tract obstruction: A microscopic

CORPORATE SOURCE: immunohistochemical study

AUTHOR: Aabe K (Reprint); Jennings R W; Harrison M R

CORPORATE SOURCE: Fukuoka Univ Hosp; Dept Pediatr Surg, Div Pediatr Surg,

AUTHOR: Matsumoto K; Perinatal Care Ctr, Jonan Ku, 7-45-1 Nankuma,

CORPORATE SOURCE: Fukuoka 8140180, Japan (Reprint); Fukuoka Univ Hosp, Dept

AUTHOR: Pediatr Surg, Div Pediatr Surg, Matern & Perinatal Care Ctr,

CORPORATE SOURCE: Jonan Ku, Fukuoka 8140180, Japan; Univ Calif San

AUTHOR: Francisco, Fetal Treatment Ctr, San Francisco, CA USA;

CORPORATE SOURCE: Univ Calif San Francisco, Dept Surg, San Francisco, CA USA

AUTHOR: abas@fukuoka-u.ac.jp

COUNTRY OF AUTHOR: ASIAN JOURNAL OF SURGERY, (JAN 2006) Vol. 29, No. 1, pp.

SOURCE: 31-35.

PUBLISHER: ISSN: 1015-9584.

LANGUAGE: ELSEVIER SINGAPORE PTE LTD, 1601, 16-F LEIGHTON CENTRE, 77

ENTRY DATE: LEIGHTON RD, CAUSEWAY BAY, HONG KONG, SAR 00000, PEOPLES R

DOCUMENT TYPE: CHINA.

LANGUAGE: Article; Journal

REFERENCE COUNT: English

ENTRY DATE: 29

AB Entered STN: 25 Jan 2007

METHODS: *ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.

RESULTS: We performed a urinary tract obstruction (UTO) surgical

CONCLUSION: procedure at 93-107 days' gestation in lambs to investigate the

CONCLUSION: relationship between pulmonary hypoplasia and the appearance of pulmonary

CONCLUSION: endocrine cells by quantitative analysis of respiratory tract cells using

CONCLUSION: light microscopic immunohistochemistry.

CONCLUSION: RESULTS: UTO produced a significant reduction in lung weight,

CONCLUSION: lung/body weight ratio, air capacity, air capacity/body weight ratio (p <

CONCLUSION: 0.01) and radial alveolar count (p < 0.05), which indicated the presence

CONCLUSION: of lung hypoplasia. These fetuses also showed a significant increase in

CONCLUSION: the number of neuron-specific enolase (NSE)-positive pulmonary endocrine

CONCLUSION: cells, expressed as the number of NSE-positive cells per bronchus (p <

CONCLUSION: 0.01) or bronchiole (p < 0.05), the number of NSE-positive cells per unit

CONCLUSION: perimeter of bronchus or bronchiole (p < 0.01), and the number of

CONCLUSION: NSE-positive cells per unit bronchial or bronchiolar surface area (p <

CONCLUSION: 0.01). CONCLUSION: These results suggest that UTO significantly retards and

CONCLUSION: modifies the structural growth and functional development of pulmonary

CONCLUSION: endocrine cells in NSE expression. We speculate that pulmonary endocrine

CONCLUSION: cells and their mediators may play a role in the problems associated with

CONCLUSION: UTO during intrauterine life.

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DOCUMENT TYPE: Article; Journal
LANGUAGE: English
REFERENCE COUNT: 27
ENTRY DATE: Entered STN: 1999

AB bombesin, which is one of the peptides produced by pulmonary neuroendocrine (PNE) cells, were carried out on the bronchioles of human congenital diaphragmatic hernia (CDH) neonates, and the findings were then compared with those in a gestational and postnatal age-matched control group. As a result, no difference was found in the number of bombesin-positive cells between the lungs of the control group and the unaffected side lungs in the CDH group except for the ratio of the bombesin-positive cells per unit of the bronchiolar surface area ($P < 0.05$). However, compared with the lungs in the control group, the affected side of the lungs in the CDH group showed a significant increase in the expression of bombesin, namely, the ratio of the bombesin-positive cells per bronchiole ($P < 0.05$), the ratio of the bombesin-positive cells per unit perimeter of the bronchioles ($P < 0.05$), and the ratio of the bombesin-positive cells per unit of the bronchiolar surface area ($P < 0.01$). These results thus suggest that hyperplasia of the PNE-cell system in the lungs of the CDH cases, especially on the affected side, exists in human fetuses. We also further speculate that PNE cells may thus play a role in the problems associated with CDH during intrauterine life in human beings.

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Corporation on STN

ACCESSION NUMBER: 1999:169812 SCISEARCH

THE GENUINE ARTICLE: 1706K

TITLE: Maternal and developmental toxicity of halogenated 4

-nitrodiphenyl ethers in mice

AUTHOR: Francis B M (Reprint); Metcalf R L; Lewis P A; Chernoff N
Univ Illinois, Dept Entomol, 1101 W Peabody Dr, Room 352,
Urbana, IL 61801 USA (Reprint); Univ Illinois, Dept
Entomol, Urbana, IL 61801 USA; US EPA, Natl Hlth &
Environm Effects Res Lab, Div Reprod Toxicol, Res Triangle
Pk, NC 27711 USA

COUNTRY OF AUTHOR: USA

SOURCE: TERATOLOGY. (FEB 1999) Vol. 59, No. 2, pp. 69-80.

ISSN: 0496-3099. JOHN WILEY & SONS INC, 605 THIRD AVE, NEW

PUBLISHER: YORK, NY 10158-0012 USA.

LANGUAGE: English; Journal

DOCUMENT TYPE: Article; Journal

REFERENCE COUNT: 42

ENTRY DATE: Entered STN: 1999

Last Updated on STN: 1999

AB In an ongoing effort to delineate structure-activity relationships in the developmental toxicity of diphenyl ethers, we evaluated the maternal and developmental toxicity of 10 diphenyl ethers related to the herbicide nitrofen. All possible trichlorophenyl 4'-nitrophenyl ethers were evaluated, as were the 2,4-difluorophenyl and 2,4-dibromophenyl 4'-nitrophenyl ethers. We also evaluated bifenoxy and chloromethoxyfen, which are 2,4-dichlorophenyl congeners with meta-substituents on the 4'-nitrophenyl ring. Nitrofen (2,4-dichlorophenyl 4'-nitrophenyl ether) was included for comparison. Identity of the halogen affected the postnatal (but not prenatal) mortality induced by 2,4-dihalogenated 4'-nitrophenyl ethers. The presence of 3'-substituents on the 4'-nitrophenyl ring reduced both pre- and postnatal toxicity of 2,4-dichlorinated congeners. Among chlorinated 4'-nitrophenyl congeners without meta-substituents on the nitrophenyl ring, the position of chlorine substituents strongly affected the congener's potential for inducing prenatal vs. postnatal syndromes. All congeners increased liver to body weight ratios in unmaternal females, but such increases were not well-correlated with either prenatal or postnatal embryotoxicity. Teratology 59:69-80, 1999, (C) 1999 Wiley-Liss, Inc.

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Corporation on STN

ACCESSION NUMBER: 1998:758559 SCISEARCH

THE GENUINE ARTICLE: 123PK

TITLE: The lungs in congenital diaphragmatic hernia: Do we

understand?

AUTHOR: Ijsestijn H; Tibboel D (Reprint)
Sophia Childrens Hosp, Dept Pediat Surg, Dr Molwaterplein
60, NL-3015 GJ Rotterdam, Netherlands (Reprint); Erasmus
Univ, Dept Pediat Surg, NL-3000 DR Rotterdam, Netherlands;
Univ Rotterdam Hosp, Sophia Childrens Hosp, Rotterdam,
Netherlands; Erasmus Univ, Dept Pediat, Div Resp Med,
Rotterdam, Netherlands

COUNTRY OF AUTHOR: Netherlands

SOURCE: PEDIATRIC PULMONOLOGY. (SEP 1998) Vol. 26, No. 3, pp.

204-218.

ISSN: 8755-6863.

PUBLISHER: WILEY-LISS, DIV JOHN WILEY & SONS INC, 605 THIRD AVE, NEW

YORK, NY 10158-0012 USA.

DOCUMENT TYPE: General Review; Journal

LANGUAGE: English

REFERENCE COUNT: 107

ENTRY DATE: Entered STN: 1998

Last Updated on STN: 1998

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Corporation on STN

ACCESSION NUMBER: 1998:681610 SCISEARCH

THE GENUINE ARTICLE: 116UT

TITLE: Congenital diaphragmatic hernia. I. Simple defect of the

diaphragm or anomaly of the pulmonary mesenchyme?

AUTHOR: Thebaud B (Reprint); de Lagausie P; Forgues D; Mercier J C

Hop Antoine Beclere, Serv Reanimat Neonatale, 157 Rue

Porte Trivaux, F-92141 Clamart, France (Reprint); Hop

Antoine Beclere, Serv Reanimat Neonatale, F-92141 Clamart,

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Lab Physiol Resp & Biol Cellulaire, F-75679 Paris, France;

Ecole Chirurg, F-75005 Paris, France

COUNTRY OF AUTHOR: France

SOURCE: ARCHIVES DE PEDIATRIE. (SEP 1998) Vol. 5, No. 9, pp.

1009-1019.

ISSN: 0929-693X.

PUBLISHER: EDITIONS SCIENTIFIQUES MEDICALES ELSEVIER, 23 RUE LINOIS.

75724 PARIS, FRANCE.

DOCUMENT TYPE: General Review; Journal

LANGUAGE: French

REFERENCE COUNT: 84

ENTRY DATE: Entered STN: 1998

Last Updated on STN: 1998

AB Described for the first time in 1846 by Bochdalek, congenital diaphragmatic hernia is still a hot topic. How can it be that a simple defect of the diaphragm still has a mortality rate reaching 50% in 1997, and this despite continuous progress in neonatal intensive care? If some problems remain unsolved, experimental studies over the past 30 years have raised some questions concerning the pathogenesis, and have shed some light into the pathophysiology of congenital diaphragmatic hernia. This article reviews the recent knowledge about the aetiology, pathogenesis and pathophysiology of this complex malformation. (C) 1998 Elsevier, Paris.

L17 ANSWER 10 OF 11 SCISEARCH COPYRIGHT (c) 2007 The Thomson

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Corporation on STN

ACCESSION NUMBER: 1997:786581 SCISEARCH

THE GENUINE ARTICLE: YC399

TITLE: Abnormal expression of pulmonary bombesin-like peptide

immunostaining cells in infants with congenital

diaphragmatic hernia

AUTHOR: Ijsestijn H (Reprint); Gaillard J L J; DeJongste J C;

Tibboel D; Cutz E

CORPORATE SOURCE: HOSP SICK CHILDREN, DEPT PATHOL, MRC, GPR LUNG DEV, TORONTO, ON M5G 1X6 CANADA; UNIV TORONTO, TORONTO, ON M5G 1X6 CANADA; ERASMUS UNIV ROTTERDAM, DEPT PEDIAT SURG, NL-3000 DR ROTTERDAM, NETHERLANDS; ERASMUS UNIV ROTTERDAM, DEPT PEDIAT, DIV RESP MED, NL-3000 DR ROTTERDAM, NETHERLANDS; ERASMUS UNIV ROTTERDAM, DEPT PATHOL, NL-3000 DR ROTTERDAM, NETHERLANDS; ERASMUS UNIV ROTTERDAM, RES INST, NL-3000 DR ROTTERDAM, NETHERLANDS; UNIV HOSP SOPHIA CHILDRENS HOSP, ROTTERDAM, NETHERLANDS; ST CLARA HOSP, ROTTERDAM, NETHERLANDS; UNIV TORONTO, TORONTO, ON M5G 1X6, CANADA; NETHERLANDS

COUNTRY OF AUTHOR: CANADA; NETHERLANDS

SOURCE: PEDIATRIC RESEARCH, (NOV 1997) Vol. 42, No. 5, pp. 715-720

ISSN: 0031-3998

PUBLISHER: WILLIAMS & WILKINS, 351 WEST CAMDEN ST, BALTIMORE, MD 21201-2436

DOCUMENT TYPE: Article; Journal

FILE SEGMENT: LIFE

LANGUAGE: English

REFERENCE COUNT: 33

ENTRY DATE: Entered STN: 1997

ABSTRACT: Last Updated on STN: 1997

*ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.

AB neonatal mortality and morbidity owing to lung hypoplasia and persistent pulmonary hypertension. Pulmonary neuroendocrine cells produce bombesin-like peptide (BLP), a peptide with growth factor-like properties involved in lung development. We examined the expression of BLP immunostaining in pulmonary neuroendocrine cells (PNEC), and in clusters of these cells called neuroepithelial bodies (NEB), in the lungs of three groups of infants: patients with CDH, newborns with lung hypoplasia due to other causes, and control subjects without lung abnormalities. Morphometric analysis included: 1) percent immunostained airways; 2) percent immunostained epithelium (i.e. frequency of PNEC and NEB); and 3) NEB size. Controls and infants with lung hypoplasia did not differ with respect to BLP immunostaining. The ipsilateral and the contralateral lungs in CDH had a similar BLP immunostaining pattern of PNEC and NEB. The BLP immunostaining varied between CDH cases, possibly due to the differences in clinical presentation. The mean NEB size was significantly increased in infants with CDH compared with the other two groups (p = 0.02). Some CDH cases with large NEBs also showed a high percentage of immunostained epithelium. Lung-body weight ratio correlated positively with percent immunostained airways, and negatively with the NEB size. We conclude that in lungs of CDH patients BLP immunostaining in PNEC and NEB differs from that of infants with lung hypoplasia due to other causes and controls. The increased BLP immunostaining observed in some cases of CDH might reflect a compensatory mechanism related to impaired lung development and/or failure of neuropeptide secretion during neonatal adaptation.

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ACCESSION NUMBER: 1997-589403 SCISEARCH

THE GENUINE ARTICLE: XP599

TITLE: Prostanoids in bronchoalveolar lavage fluid do not predict outcome in congenital diaphragmatic hernia patients

AUTHOR: Ijsselstijn H (Reprint); Zijlstra F J, deJongste J C, Tibboel D

CORPORATE SOURCE: ERASMUS UNIV ROTTERDAM, DEPT PAEDIAT SURG, DIV RESP MED, ROTTERDAM, NETHERLANDS; ERASMUS UNIV ROTTERDAM, DEPT PAEDIAT, DIV RESP MED, ROTTERDAM, NETHERLANDS; ERASMUS UNIV ROTTERDAM, DEPT PHARMACOL, ROTTERDAM, NETHERLANDS; UNIV ROTTERDAM HOSP, SOPHIA CHILDRENS HOSP, ROTTERDAM, NETHERLANDS

COUNTRY OF AUTHOR: NETHERLANDS

SOURCE: MEDIATORS OF INFLAMMATION, (JUN 1997) Vol. 6, No. 3, pp. 217-224

ISSN: 0962-9351

PUBLISHER: HINDAWI PUBLISHING CORPORATION, PO BOX 1210, SYLVANIA, OH 43560 USA.

DOCUMENT TYPE: Article; Journal

LANGUAGE: English

REFERENCE COUNT: 1

ENTRY DATE: Last Updated on STN: 15 Sep 2005

*ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS.

AB Vasoactive prostanoids may be involved in persistent pulmonary hypertension (PPH) in infants with a congenital diaphragmatic hernia (CDH). We hypothesized that increased levels of prostanoids in bronchoalveolar lavage (BAL) fluid would predict clinical outcome. We measured the concentrations of 6-keto-prostaglandin F-1 alpha (6-keto-PGF(1 alpha)), thromboxane B-2 (TxB(2)), protein, albumin, total cell count, and elastase-alpha(1)-proteinase-inhibitor complex in BAL fluid of 18 CDH patients and of 13 control subjects without PPH. We found different concentrations of prostanoids in BAL fluid of CDH patients with PPH: infants with a poor prognosis had either high levels of both 6-keto-PGF(1 alpha) and TxB(2) compared to controls, or high levels of 6-keto-PGF(1 alpha) only. TxB(2) levels showed a large variability in all CDH patients irrespective of outcome. We conclude that prostanoid levels in BAL fluid do not predict clinical outcome in CDH patients.

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FILE 'CAPLUS' ENTERED AT 20:32:20 ON 12 APR 2007

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FILE 'CAPLUS' ENTERED AT 20:32:43 ON 12 APR 2007

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FILE 'SCISEARCH' ENTERED AT 20:50:35 ON 12 APR 2007

L10 7 S E2

L11 727 S TSUTSUMI Y2/AU

L12 6 S L11 AND CALCITONIN

L13 19 S E3

L14 14 S L13 NOT L6

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L16 12 S E4

L17 11 S L16 NOT (L15 OR L10 OR L6)

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